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### TIME RESOLVED PHOTOLUMINESCENCE SPECTRA OF A MID-INFRARED MULTIPLE QUANTUM WELL SEMICONDUCTOR LASER

THESIS

Anthony L. Franz, Captain, USAF

AFIT/GAP/ENP/97D-04

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### THESIS

Presented to the Faculty of the Graduate School of Engineering of the

Air Force Institute of Technology

Air University

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In Partial Fulfillment of the Requirements for the

Degree of Master of Science in Engineering Physics

Anthony L. Franz

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December, 1997

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### Abstract

Recombination mechanisms in mid-IR semiconductor lasers are strongly dependent on the carrier density of the active region. Therefore, in order to determine the importance of different recombination mechanisms, an accurate knowledge of the carrier density is important. The objective of this research is to improve previous carrier density estimates through the incorporation of spectral information. In the long run, this will help researchers to more accurately characterize the recombination mechanisms and improve the efficiency of the lasers at these wavelengths.

One hundred photoluminescence (PL) spectra were calculated for a variety of carrier densities. Calculations were made for an InAsSb/InAlAsSb multiple quantum well laser sample assuming parabolic bands. In order to compare the results with experiment, the calculated spectra were convolved with the measured pump beam spectrum to account for broadening in the measurement process. The widths of the convolved calculated spectral profiles were tabulated as a function of carrier density.

Actual spectra were measured using the Ultrafast Mid-Infrared Photoluminesce System, which uses upconversion to measure the PL intensity in time steps smaller than 1 ps. PL spectra were obtained at 30 times, ranging from 100 ps to 3 ns. Spectral widths were measured and tabulated as a function of time.

Combining the plot of convolved calculated spectral width vs. carrier density with the plot of measured spectral width vs. time, we were able to describe the variation of carrier density with time. The carrier density vs. time plot thus generated agreed with

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earlier measurements by Cooley for low carrier densities. The discrepancy at higher carrier densities could be due to changing experimental conditions or the break down of the parabolic band approximation at higher carrier densities.

## TIME RESOLVED PHOTOLUMINESCENCE SPECTRA OF A MID-INFRARED MULTIPLE QUANTUM WELL SEMICONDUCTOR LASER

### 1. Introduction

### Background

Mid-infrared semiconductor lasers are being developed for infrared (IR) countermeasures and chemical analysis. These lasers take advantage of the 2-5  $\mu$ m transparency window in the atmosphere and have applications in remote sensing and environmental monitoring. Past research has analyzed the nonradiative recombination mechanisms in these structures in an attempt to minimize them. Auger recombination can be a major contributor to loss of efficiency in these devices resulting in increased lasing threshold currents at higher temperatures (Agrawal, 1993: 98-100). Techniques to study the carrier dynamics of these devices will help find solutions to the Auger problem.

Capt William Cooley constructed the Ultrafast Mid-Infrared Photoluminescence System (UMIPS) as part of his dissertation research. UMIPS uses sum frequency generation (SFG), or upconversion, to examine the photoluminescence (PL) lifetime of lasers emitting in the mid-IR (Cooley, 1996). We have extended his technique to measure PL spectra at 0.1 picosecond intervals. Analysis of the resulting spectra can provide insight into the carrier density at each time the PL is measured.

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After the sample is optically pumped with a pulse the carrier density will decrease over time as electrons and holes recombine. The typical model used to describe this decrease for situations below threshold is a rate equation

$$-\frac{\partial n}{\partial t} = An + Bn^2 + Cn^3 \tag{1}$$

where A, B, and C are the Shockley-Read-Hall, radiative, and Auger coefficients, respectively and n is the carrier density (Agrawal, 1993: 38). It is assumed that the number of electrons equals the number of holes. Each coefficient describes a type of recombination where the power of n in each term depends on the number of electrons and holes required for the type of recombination.

The types of recombination considered are illustrated in Figure 1. Shockley-Read-Hall recombination occurs when a defect is present in the lattice. This defect has a localized continuous density of states and the electron returns to the valence band through these states (Agrawal, 1993: 119). Radiative recombination occurs when an electron transitions to an empty state (a hole) in the valence band. The excess energy is released as a photon which has a frequency equal to the energy difference between the two states. Band-to-band Auger recombination occurs when the excess energy from a downward transition is used to shift another electron into a higher energy state (Agrawal, 1993: 99). In direct gap semiconductors there are three types of Auger recombination processes which are differentiated by the bands involved (Agrawal, 1993: 99). These processes are shown in Figure 2. We are primarily interested in the CCCH Auger process.

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### Approach

This effort used calculated and measured spectral data to find an estimate of the carrier density as a function of time. PL spectra were calculated based on a given carrier density. As the number of available carriers and holes decrease, fewer energy levels in the quantum wells will be occupied and the width of the PL spectra becomes narrower. The peak may also shift to slightly lower energies, depending on the range of carrier densities observed, as more radiative recombination occurs between lower energy levels. These calculated spectra had to be corrected for broadening that occurs in the experiment. To correct the calculated data for this effect, the spectrum of the upconversion pump beam used in the experiment was measured and convolved with the calculated PL spectra. The widths of the resulting convolved calculated spectra were obtained as a function of carrier density as illustrated in Figure 3a.

The measured spectral data were provided by Capt Craig Largent of the Engineering Physics Department at AFIT. He used the UMIPS to measure time-resolved spectra from the sample at different delay times after the excitation of the sample. The widths of these measured PL spectra were obtained as a function of time as illustrated in Figure 3b.

In order to determine carrier density as a function of time, the measured widths and convolved calculated widths were compared. This process is illustrated in Figure 3. Each delay time has a measured width associated with it. This measured width can be compared to the convolved calculated widths to find the one that matches it. The matching convolved calculated width is associated with a carrier density. This process

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Figure 3: Schematic showing how carrier density was linked to time. The plots are illustrative and do not represent real data. (a) is a plot of the convolved calculated spectral widths vs. carrier density, and (b) is a plot of the measured widths vs. time. First a width  $w_1$  is found for a given time  $t_1$  using (b). Then  $w_1$  is associated with a carrier density  $n_1$  using (a). The result of this is a pair of values  $(t_1, n_1)$ .

was repeated for each time to find the carrier density as a function of time. This relation was then plotted and compared to Cooley's results.

### **Overview**

Chapter 2 will address the theoretical calculation of PL spectra for different carrier densities. Chapter 3 will discuss the UMIPS and the measured data obtained by Capt Largent. Chapter 4 will present the results of the spectral width calculation and discuss the convolution of the calculated spectra with the measured pump beam. The carrier density as a function of time results and conclusions from the analysis will be presented. Chapter 5 is a summary of the research and recommendations for future research in this area.

### 2. Theory

### **Overview**

This chapter will address the theoretical aspects of the research. The goal of the theoretical calculations is to determine the width of the PL spectra that are expected for different carrier densities. The first step is to calculate the energy levels in the quantum wells. This is the topic of the second section. The third section discusses how the spontaneous emission spectra are calculated, and the fourth section discusses the convolution of these spectra performed to account for intraband relaxation. The last section is a summary of the calculation process.

It should be noted that a key assumption made in the calculations is that the band structure of the sample is parabolic so that

$$E_{e}(k_{e}) = E_{c} + \frac{\hbar^{2}k_{e}^{2}}{2m_{e}}, \quad E_{h}(k_{h}) = E_{v} - \frac{\hbar^{2}k_{h}^{2}}{2m_{h}}$$
(2)

where  $E_e$  and  $E_h$  are the energies of an electron and hole,  $k_e$  and  $k_h$  are the wave vectors for the electron and hole,  $E_c$  and  $E_v$  are the conduction and valence band edges, and  $m_e$ and  $m_h$  are the effective masses of the electrons and holes. The quality of the calculated spectra will depend on the quality of this assumption.

#### **Energy Levels in the Quantum Wells**

The first step in the calculation is to find the energy levels for electrons in the conduction band and for heavy and light holes in the valence band. The model used to calculate the energy levels is a single finite potential well. This is a standard problem in quantum mechanics texts and the results are well documented (Liboff, 1989: 256-65;



Figure 4: Finite square well model of the quantum well used in energy level calculations.

Cohen-Tannoudji, 1977: 74-7). In this model the potential energy surface is modeled as a square well in the z direction as shown in Figure 4. This model assumes a parabolic band structure and, by applying it to the multiple quantum well structure, that each well is isolated from neighboring wells.

The energy levels must be calculated for each band separately. Using the envelope functions as the wavefunctions and setting the electron momentum to zero the Schrödinger equation becomes

$$-\frac{\hbar^2}{2m}\frac{d^2}{dz^2} + VF_z = EF_z \tag{3}$$

where *m* is the effective mass, *V* is the potential,  $F_z$  is the envelope function in the *z* direction, and *E* is the energy of the band. It should be noted that both *m* and *V* have different values in the well and barrier regions. The effective mass for electrons, heavy holes, or light holes in each region is used when calculating the energy levels for the conduction, heavy hole, or light hole band respectively. Also, for reasons to be described

later, the value for  $V_0$  in Figure 4 will be different for the conduction, heavy hole, and light hole bands. Plane wave solutions are assumed for the envelope functions and the following boundary conditions are applied at the interfaces (Corzine, 1993: 59-60):

$$F_{z,well} = F_{z,barrier}$$
 and  $\frac{1}{m_{well}} \frac{dF_{z,well}}{dz} = \frac{1}{m_{barrier}} \frac{dF_{z,barrier}}{dz}$  (4)

There are two classes of solutions for the envelope functions: even and odd. The even solutions are cosines in the well and decay exponentially in the barrier; the odd solutions are sines in the well and also decay exponentially in the barrier (Liboff, 1989: 260). Using  $k_z^2 = 2m_{well}E/\hbar^2$  and  $\alpha_z^2 = (2m_{barrier}/\hbar^2)(V_0 - E)$ , the characteristic equations for

energy for even and odd solutions are

$$\tan\left(k_{z}\frac{L_{z}}{2}\right) = \frac{m_{well}}{m_{barrier}}\frac{\alpha_{z}}{k_{z}} \qquad \text{for even solutions}$$

$$\cot\left(k_{z}\frac{L_{z}}{2}\right) = -\frac{m_{well}}{m_{barrier}}\frac{\alpha_{z}}{k_{z}} \qquad \text{for odd solutions} \qquad (5)$$

where  $L_z$  is the width of the quantum well and  $V_0$  is the depth of the well. These characteristic equations can only be solved for discrete values of energy,  $E_n$  (Corzine, 1993a: 60-1). The energy levels for the conduction, heavy hole, and light hole bands,  $E_{cn}$ ,  $E_{hhn}$ , and  $E_{lhn}$  respectively, are determined in this manner using the appropriate effective masses and well depths.

The depth of the well is determined by the energy gaps of the well and barrier and the band offsets. The offsets are a property of the material, but several factors can change the gap for a given material. A major factor is the temperature of the system. The temperature dependence will be assumed to follow the Varshni relation

$$E_g(T) = E_g(0) - \frac{\alpha T^2}{\beta + T}$$
(6)

where  $E_g(0)$  is the gap at T = 0 and  $\alpha$  (in eV/K) and  $\beta$  (in K) are constant parameters fit to experimental data (Varshni, 1967: 149).

In addition to temperature, the energy gap for the well will be affected by strain. Stress in quantum well structures occurs when the lattice parameter of the well material is different than the lattice parameter of the substrate and the well layer is thinner than a critical thickness. If the well is thicker than the critical thickness then the lattice will crack to relieve the strain. If the well is thinner then the lattice will not crack and will be under compressive or tensile strain as shown in Figure 5 (O'Reilly, 1989:122-4). Figure 6 shows the effect of compressive strain on the band structure of the well. There is little affect on the conduction band, but the valence band changes drastically. The gap between the conduction band and valence band increases and the degeneracy of the heavy and light holes is removed. Assuming parabolic bands at zone center, for III-V compound semiconductors under a strain given by

$$\varepsilon_s = \frac{a_{0, \text{ substrate}} - a_{0, \text{ well}}}{a_{0, \text{ well}}}$$
(7)

the energy band shifts are given by

$$\Delta E_{hh} = -2a\varepsilon_s \frac{C_{11} - C_{12}}{C_{11}} + b\varepsilon_s \frac{C_{11} + C_{12}}{C_{11}}$$

$$\Delta E_{lh} = -2a\varepsilon_s \frac{C_{11} - C_{12}}{C_{11}} - b\varepsilon_s \frac{C_{11} + 2C_{12}}{C_{11}}$$
(8)



Figure 5: Strain in semiconductor layers. (a) Three unstrained layers where the middle layer has a larger lattice parameter. (b) The middle layer distorts and is under compressive strain when placed between the two outer layers.



where a is the hydrostatic deformation potential, b is the shear deformation potential, and the Cs are elastic stiffness coefficients (Coleman, 1993: 377). For a well under compressive strain the energy gap is given by  $E_g(T) + \Delta E_{hh}$ .

Thus when strain is present and the conduction band offset does not equal  $\frac{1}{2}$ , three separate well depths must be considered: one each for electrons (conduction band), heavy holes, and light holes. Equation (5) must be applied to each band with the appropriate effective masses and well depth,  $V_0$ . For a layer under compressive strain the well depths are given by

$$V_{cb} = \Delta E_{cb} \left[ E_{g,barrier} \left( T \right) - E_{g,well} \left( T \right) - \Delta E_{hh} \right]$$

$$V_{hh} = \left[ 1 - \Delta E_{cb} \right] \left[ E_{g,barrier} \left( T \right) - E_{g,well} \left( T \right) - \Delta E_{hh} \right]$$

$$V_{lh} = \left[ 1 - \Delta E_{cb} \right] \left[ E_{g,barrier} \left( T \right) - E_{g,well} \left( T \right) - \Delta E_{hh} \right] - \Delta E_{lh}$$
(9)

where  $\Delta E_{cb}$  is the conduction band offset expressed as a fraction of one. A comparison of a strained and an unstrained well is shown in Figure 7.



### **Spontaneous Emission Spectrum**

The total spontaneous emission rate per unit volume over a given energy range is found by multiplying the transition rate per unit volume times the number of optical modes in the energy range:

$$R_{sp}(\hbar\omega) d(\hbar\omega) = W_{c \to v} \left[ V \rho_{opt}(\hbar\omega) d(\hbar\omega) \right]$$
(10)

where  $W_{c \to v}$  is the transition rate from the conduction band to the valence band, V is the volume of the active region, and  $\rho_{opt}$  is the optical density of states (Corzine, 1993:42).

The transition rate is found using Fermi's Golden Rule, which is derived using time-dependent perturbation theory to solve Schrödinger's equation when the Hamiltonian has the form  $H = H_0 + \lambda H'(t)$ . In general the transition rate, W, from state a to state b is given by

$$W_{a \to b} = \frac{2\pi}{\hbar} \left| H'_{ab} \right|^2 \rho(E) \tag{11}$$

where  $H'_{ab} = \langle b|H'|a \rangle$ , H' is the perturbation term in the Hamiltonian and  $\rho(E)$  is the density of states as a function of energy (Bransden, 1983: 111-6 and Liboff, 1989: 576-84). Spontaneous emission involves the interaction of the electron in the conduction band with an electromagnetic wave. The wave interaction term is treated as a perturbation to the system so that

$$H'(\mathbf{r}) \equiv \frac{e}{2m_0} A(\mathbf{r}) \hat{\mathbf{e}} \cdot \mathbf{p} \text{ and } H'_{eh} \equiv \left\langle \psi_h \middle| H'(\mathbf{r}) \middle| \psi_e \right\rangle$$
(12)

where e and  $m_0$  are the charge and mass of an electron, A is the vector potential of the spontaneous emission wave,  $\hat{\mathbf{e}}$  is the polarization unit vector of the wave, and  $\mathbf{p}$  is the

momentum operator (Corzine, 1993: 28-9).  $\psi_i$  are the wavefunctions of the electrons and holes and are solutions to the Schrödinger equation when  $H = H_0$ . Using the envelope function approximation,

$$\psi_i \equiv F_i(\mathbf{r}) \, u_i(\mathbf{r}) \tag{13}$$

where  $F_i$  is the envelope function and  $u_i$  is the Bloch function. The key assumption is that the Bloch functions are not a strong function of k (Corzine, 1993: 19). The Bloch functions are periodic with the crystal lattice (McKelvey, 1993: 319) and the envelope functions are normalized plane waves for unconfined directions in the crystal. The envelope functions for the confined direction of the well were discussed on page 8 (Corzine, 1993: 20,59).

For momentum to be conserved in the transition,  $\mathbf{k}_{h} = \mathbf{k}_{e} + \mathbf{k}_{photon}$  must be valid. Typically the wavelength of the light is much greater than the De Broglie wavelength of the electron. If this is true then  $\mathbf{k}_{h} = \mathbf{k}_{e}$  and the transitions are said to obey **k**-selection rules. This applies to band-to-band transitions, not localized transitions, where the envelope functions of the initial and final states are plane waves (Corzine, 1993: 30).

For a quantum well the electrons and holes are confined along the growth direction so the z direction envelope functions must be included in the wavefunctions. If the wavelength of the light is much larger than the width of the well then the vector potential is can be considered to be a constant,  $A_0$ , in the region of the well. Using these assumptions and substituting Equation (13) into Equation (12) gives

$$\left|H_{eh}'\right|^{2} = \left(\frac{eA_{0}}{2m_{0}}\right)^{2} \left|M_{T}\right|^{2}, \quad \left|M_{T}\right|^{2} \equiv \left|\left\langle u_{v}\left|\hat{\mathbf{e}}\cdot\mathbf{p}\right| u_{c}\right\rangle\right|^{2} \left|\left\langle F_{h}\left|F_{e}\right\rangle\right|^{2}$$
(14)

where  $|M_T|^2$  are the transition matrix elements which will be described below (Corzine, 1993: 30-1).

Equation (14) is substituted into Fermi's Golden Rule (Equation (11)). The transition rate per unit volume can be expressed as

$$W_{c \to v} = \frac{2\pi}{\hbar} \left( \frac{eA_0}{2m_0} \right)^2 \left| M_T \right|^2 \rho_{red} \left( E_{eh} - E'_g \right) f_c \left( 1 - f_v \right)$$
(15)

where  $\rho_{red}$  is the reduced density of states,  $E_{eh}$  is the transition energy  $\hbar\omega$ ,  $E'_g$  is the gap between the subbands of the transition, and  $f_c$  and  $f_v$  are the Fermi-Dirac distributions for electrons in the conduction and valence bands respectively (Corzine, 1993; 33-5). These terms are discussed individually in the following sections.

### Density of States

Fermi's Golden Rule has the density of states as a factor. The density of states for a quantum well is different than for a bulk semiconductor. A bulk crystal with dimensions  $L_x$ ,  $L_y$ , and  $L_z$  is modeled as a square well with infinitely high walls in each dimension (the electrons are confined to the crystal). Requiring the wavefunctions to be zero at the boundaries gives standing waves for the wavefunctions of the form

$$\varphi(x,y,z) = \left(\frac{8}{L_x L_y L_z}\right)^{\frac{1}{2}} \sin\left(\frac{n_x \pi x}{L_x}\right) \sin\left(\frac{n_y \pi y}{L_y}\right) \sin\left(\frac{n_z \pi z}{L_z}\right)$$
(16)

where  $n_i$  are the quantum numbers in each direction (Cohen-Tannoudji, 1977: 199).

The wave vector has the form  $\mathbf{k} = k_x + k_y + k_z$  where  $k_i = n_i \pi/L_i$ . The k vector sweeps out a sphere of volume  $V_k = \frac{4}{3} \pi k^3$  which contains a number of states,  $N_s$ , given by

$$N_{s} = V_{k} \frac{L_{x} L_{y} L_{z}}{(2\pi)^{3}}$$
(17)

where spin has been neglected. Spin will be addressed with the transition matrix elements. In general, if  $\rho(k)$  is the density of states then

$$\int \rho(k)dk = N_s/V \tag{18}$$

when integrating over  $V_k$ , so that

$$\rho(k) = \frac{1}{V} \frac{dN_s}{dk} \tag{19}$$

where V is the total volume of the crystal (Corzine, 1993: 23-4). We are interested in the energy of the electron, not its wave vector. The density of states as a function of energy can be found through the relation

$$\rho(E)dE = \rho(k)dk \to \rho(E) = \frac{\rho(k)}{dE/dk}$$
(20)

which requires a knowledge of how the energy depends on the wave vector (Corzine, 1993: 24). For the parabolic band approximation dE/dk can be found easily from Equation (2):

$$E(k) = \frac{\hbar^2 k^2}{2m} \rightarrow \frac{dE}{dk} = \frac{\hbar^2}{m} k$$
(21)

This relationship applies to the conduction and valence bands when the appropriate effective masses are used.

For a bulk semiconductor  $V_k$  is a sphere as shown in Figure 8 and the density of states is found by plugging Equation (17) into Equation (19)



semiconductor. (b) k space for a quantum well semiconductor. The levels along the z (growth) direction have a larger spacing than the in-plane directions (after Figure 1 in Corzine, 1993: 22).

$$\rho(k) = \frac{1}{V} \frac{dN_s}{dk} = \frac{1}{V} \frac{d}{dk} \left( \frac{\frac{4}{3}\pi k^3}{(2\pi)^3 / (L_x L_y L_z)} \right) = \frac{1}{(2\pi)^3} \frac{L_x L_y L_z}{V} \frac{4}{3} 3\pi k^2 = \frac{k^2}{2\pi^2}$$
(22)

Substituting Equations (21) and (22) into Equation (20) gives

$$\rho(E) = \frac{\sqrt{E}}{4\pi^2} \left(\frac{2m}{\hbar^2}\right)^{\frac{N}{2}}$$
(23)

for a bulk semiconductor with parabolic bands, ignoring spin degeneracy (Corzine, 1993: 23-5).

For a quantum well with  $L_z \ll L_x$  and  $L_y$ ,  $k_z \gg k_x$  and  $k_y$  and the distance between consecutive levels of  $k_z$  will be very large compared to the spacing between consecutive levels of  $k_x$  and  $k_y$  as shown in Figure 8. The k vector will sweep out circles in each  $k_z$  level so that  $V_k = \pi k^2$  and one state occupies an area of  $(2\pi)^2/(L_x L_y)$ . N<sub>s</sub> will equal  $V_k$  divided by this area so the density of states is

$$\rho(k) = \frac{1}{V} \frac{dN_s}{dk} = \frac{1}{V} \frac{d}{dk} \left( \frac{2\pi k^2}{(2\pi)^2 / (L_x L_y)} \right) = \frac{1}{(2\pi)^2} \frac{L_x L_y}{V} 2\pi k = \frac{k}{2\pi L_z}$$
(24)

for a quantum well. Substituting Equations (21) and (24) into Equation (20) gives

$$\rho(E) = \frac{m}{2\pi\hbar^2 L_z} \tag{25}$$

for each plane of k states in a two dimensional quantum well semiconductor with parabolic bands, ignoring spin degeneracy. This form of the density of states is a step function with the step size given in Equation (25). A new step occurs at each energy level of the quantum well (Corzine, 1993: 23-5). The density of states functions for a bulk (3D) and quantum well (2D) semiconductor are compared in Figure 9.



Figure 9: Comparison of the density of states,  $\rho(E)$ , for a quantum well and a bulk semiconductor. As the width of the quantum well increases the step size decreases and the quantum well case will approach the bulk case (after Figure 2 in Corzine, 1993: 25).

The reduced density of states is defined as (Corzine, 1993: 34)

$$\rho_{red} \equiv \frac{\rho(k)}{dE_{eh}/dk}$$
(26)

where  $E_{eh}$  is the transition energy,  $\hbar\omega$ , given by

$$E_{eh} \equiv E_e - E_h = E'_g + \frac{\hbar^2 k^2}{2m_c} + \frac{\hbar^2 k^2}{2m_v}$$
(27)

for the parabolic bands, where  $E'_g$  is the gap between the subbands of the transition. Taking the derivative of this results in

$$\frac{dE_{eh}}{dk} = \frac{\hbar^2 k}{m_r}$$
, where  $\frac{1}{m_r} = \frac{1}{m_c} + \frac{1}{m_v}$  (28)

with  $m_r$  defined as the reduced mass. The reduced density of states,  $\rho_{red}$ , has the form of Equation (25) with the mass replaced by the reduced mass and E by  $E_{eh}$  -  $E_{g}$  (Corzine, 1993: 34).

### Quasi-Fermi Functions

The transition rate given by Fermi's Golden Rule is for a transition from one state in the conduction band to one state in the valence band. The possibility that some states may be occupied must be taken into account. The transition rate is multiplied by the probability of having an electron present in the conduction band state and not having an electron present in the corresponding valence band state (presence of having a hole).

When the laser impinges on the sample, electrons from the valence band are excited into the conduction band. Intraband relaxation occurs on the order of picoseconds while recombination takes tens of picoseconds or more. After intraband relaxation and before the start of recombination, the electrons occupy the lowest energy states available



according to the Fermi-Dirac distribution with an appropriate quasi-Fermi level. This quasi-equilibrium condition is illustrated in Figure 10. The Fermi-Dirac distribution with a corresponding quasi-Fermi level is

$$f_{c} = \left[1 + \exp\left(\frac{E_{e} - E_{Fc}}{k_{B}T}\right)\right]^{-1} \text{ for the conduction band}$$

$$f_{v} = \left[1 + \exp\left(\frac{E_{h} - E_{Fv}}{k_{B}T}\right)\right]^{-1} \text{ for the valence band}$$
(29)

where  $E_e$  is the electron energy,  $E_h$  is the hole energy,  $k_B$  is Boltzmann's constant, and T is the temperature of the carriers. The quasi-Fermi levels,  $E_{Fc}$  and  $E_{Fv}$ , are the energies at which the probability of an electron occupying a state in the band is  $\frac{1}{2}$  (Verdeyen, 1995: 450 and Kittel, 1980: 379). The total carrier density, N for the conduction band and P for the valence band, can be calculated by integrating the product of the density of states (times two to account for spin) and the quasi-Fermi function over the entire band. For parabolic bands this is

$$N = 2 \int_{E_c}^{\infty} \rho^{2D} \left( E - E_c \right) f_c dE$$

$$P = 2 \int_{-\infty}^{E_v} \rho^{2D} \left( E_v - E \right) \left( 1 - f_v \right) dE$$
(30)

where  $E_c$  and  $E_v$  are the conduction and valence band edges and  $\rho^{2D}$  is the two dimensional density of states. Plugging Equations (25) and (29) into Equation (30) and integrating gives, for the conduction band,

$$N = \frac{m_c k_B T}{\pi \hbar^2 L_z} \sum_n \ln \left[ 1 + \exp\left(-\frac{E_{cn} - E_{Fc}}{k_B T}\right) \right]$$
(31)

where the summation is over the energy levels in the conduction band. For a given total carrier density a quasi-Fermi level can be calculated numerically (Corzine, 1993: 44). The quasi-Fermi level for the valence band is found by replacing N with P and summing over the energy levels in the heavy and light hole subbands.

### Transition Matrix Elements

The transition matrix elements are defined in Equation (14). The envelope function term will be close to unity when the conduction band and the valence band state have the same quantum numbers and close to zero when they have different quantum numbers (Corzine, 1993: 31-2). It is assumed that  $|\langle F_h | F_e \rangle|^2$  is unity for these allowed transitions  $(n_c = n_v)$  and zero for other transitions  $(n_c \neq n_v)$ . This reduces  $|M_T|^2$  to

$$\left|M_{T}\right|^{2} = \left|\left\langle u_{v} \mid \hat{\mathbf{e}} \cdot \mathbf{p} \mid u_{c}\right\rangle\right|^{2}$$
(32)

and the Bloch functions must be now be considered.

Semiconductor band structure is frequently modeled as shown in Figure 11. These bands can be thought of as originating from atomic orbitals where the conduction band corresponds to as s orbital and the valence bands correspond to the  $p_x$ ,  $p_y$ , and  $p_z$  orbitals (McKelvey, 1993: 365-6). The Bloch functions for these orbitals are labeled  $u_s$ ,  $u_x$ ,  $u_y$ , and  $u_z$ , respectively, and have the same symmetry properties of the corresponding atomic orbitals. These symmetry relations combined with the momentum operator give the following relations:

$$\left\langle \begin{array}{c} u_{s} \left| p_{i} \right| u_{j} \right\rangle = 0, \text{ for } i \neq j \\ \left\langle \begin{array}{c} u_{s} \left| \mathbf{p} \right| u_{i} \right\rangle = \left\langle \begin{array}{c} u_{s} \right| p_{i} \right| u_{i} \right\rangle \equiv M \\ \left\langle \begin{array}{c} u_{s} \left| \mathbf{p} \right| \left| \overline{u}_{i} \right\rangle = 0 \end{array} \right.$$

$$(33)$$



Figure 11: Subband structure for an unstrained semiconductor. The subbands are parabolic as represented by each bands' effective mass.  $E_g$  is the energy gap between the conduction and valence bands.  $\Delta_{so}$  is the gap between the spin-orbit subband and the heavy and light hole subbands (after Figure 7 in Corzine, 1993: 49)

where i = x, y, z and  $u_i$ ,  $\overline{u}_i$  indicate spin-up and spin-down functions (Corzine, 1993: 45-6). *M* is the orbital momentum matrix element and is a property of the material. The valence band Bloch functions  $u_{hh}$ ,  $u_{ih}$ , and  $u_{so}$  can be written as linear combinations of the orbital Bloch functions. For electron **k** vectors directed along the *z* direction

$$u_{hh} = -\frac{1}{\sqrt{2}} \left( u_x + iu_y \right), \qquad \overline{u}_{hh} = \frac{1}{\sqrt{2}} \left( \overline{u}_x + i\overline{u}_y \right)$$

$$u_{lh} = -\frac{1}{\sqrt{6}} \left( \overline{u}_x + i\overline{u}_y - 2u_z \right), \quad \overline{u}_{lh} = \frac{1}{\sqrt{6}} \left( u_x - iu_y + 2\overline{u}_z \right)$$

$$u_{so} = -\frac{1}{\sqrt{3}} \left( \overline{u}_x + i\overline{u}_y + u_z \right), \quad \overline{u}_{so} = \frac{1}{\sqrt{3}} \left( u_x - iu_y - \overline{u}_z \right)$$
(34)

where six equations are used to account for the spin-up and spin-down states (Corzine, 1993: 47). Including these spin states there are four possible transitions from the conduction band to each valence band. Setting the overlap integral to unity as described on page 20 gives

$$\left|M_{T}\right|_{v}^{2} = \sum_{u_{c}, \bar{u}_{c}} \sum_{u_{v}, \bar{u}_{v}} \left|\left\langle u_{v} \middle| \hat{\mathbf{e}} \cdot \mathbf{p} \middle| u_{c} \right\rangle\right|^{2}$$
(35)

Expanding the dot product into its components, replacing  $u_v$  in Equation (35) with the appropriate relations for each band from (34) and using the selection rules in (33) gives the relative transition strength with respect to  $|M|^2$  for transitions to each valence band as

$$\frac{\left|M_{T}\right|_{\nu}^{2}}{\left|M\right|^{2}} = \begin{cases} 1 - \left|\hat{\mathbf{k}} \cdot \hat{\mathbf{e}}\right|^{2} & \text{for the heavy hole band} \\ \frac{1}{3} + \left|\hat{\mathbf{k}} \cdot \hat{\mathbf{e}}\right|^{2} & \text{for the light hole band} \\ \frac{2}{3} & \text{for the spin - orbit band} \end{cases}$$
(36)

where  $\hat{\mathbf{k}}$  is the unit vector of the electron wave vector which was set to the *z* direction (Corzine, 1993: 50-1).



subbands. The strengths labeled are in units of  $|M|^2$  (after Figure 9 in Corzine, 1993; 53).

By averaging over all of the possible directions of the k vector in the first quadrant of k space it is seen that for a quantum well the average direction is the growth or z direction. Because of the dot product in Equation (36), the transition strength is a function of the angle between  $\hat{k}$  and  $\hat{e}$ . This angular dependence is plotted in Figure 12 for the heavy and light hole subbands. Setting  $\hat{k}$  equal to  $\hat{k}_{ave}$  allows the transition strengths to be found for three orthogonal  $\hat{e}$  set in directions relative to the quantum well as shown in Figure 13 (Corzine, 1993: 52-5). We are interested in all spontaneous emission regardless of polarization so the  $|M_T|^2$  term in Equation (15) needs to be replaced with an average over the three polarization directions,  $|M_{ave}|^2$ , defined as (Corzine, 1993: 43)


$$\left|M_{ave}\right|^{2} \equiv \frac{1}{3} \sum_{\substack{\text{all three} \\ \text{polarizations}}} \left|M_{T}\right|^{2}$$
(37)

For the transitions in a quantum well, substituting Equation (36) into Equation (37) gives

$$\left|M_{ave}\right|^{2} = \begin{cases} \frac{1}{3}(0+1+1)|M|^{2} \\ \frac{1}{3}(\frac{4}{3}+\frac{1}{3}+\frac{1}{3})|M|^{2} \\ \frac{1}{3}(\frac{2}{3}+\frac{2}{3}+\frac{2}{3})|M|^{2} \end{cases} = \frac{2}{3}|M|^{2}$$
(38)

for transitions to all bands. The values for the heavy hole and light hole transitions came from Figure 13. Equation (38) is used in Equation (15) in place of the  $|M_T|^2$  term.

 $|M|^2$  must be determined for the sample to quantify the spontaneous emission rate. Using the **k**·**p** technique for the four bands in Figure 11 an equation for the approximate conduction band effective mass,  $m^*$ , can be rearranged to solve for  $|M|^2$ (Corzine, 1993: 48-9)

(39)

$$|M|^{2} = \left(\frac{m_{0}}{m^{*}} - 1\right) \frac{E_{g} + \Delta_{so}}{2\left(E_{g} + \frac{2}{3}\Delta_{so}\right)} m_{0}E_{g}$$

It should be noted that  $m^*$  is not the true effective mass because contributions from higher and lower level bands will make the true effective mass heavier and lighter respectively. The inaccuracy in Equation (39) depends on how large an effect these unaccounted for bands have on the true effective mass (Corzine, 1993: 48-9 and Yan, 1990: 214-5).

## **Optical Density of States**

The optical density of states in an energy range is found by using the 3D density of states,  $\rho(k)$ , found in Equation (22). For photons

$$k_{opt} = n_r \frac{2\pi}{\lambda} = n_r \frac{\omega}{c} = \frac{n_r}{\hbar c} (\hbar \omega), \quad \frac{dk_{opt}}{d(\hbar \omega)} = \frac{n_g}{\hbar c}$$
(40)

where  $n_r$  and  $n_g$  are the index of refraction and the group index of refraction. Substituting these relations into Equation (20) yields

$$\rho_{opt}(\hbar\omega) = \frac{1}{\pi^2} \frac{n_r n_g}{(\hbar c)^3} (\hbar\omega)^2$$
(41)

in units of energy<sup>-1</sup> cm<sup>-3</sup>, where a factor of two has been multiplied to account for the two polarization states (Corzine, 1993: 42).

## Vector Potential

The vector potential is found by relating the energy in the field to the energy of a photon. Solving for the vector potential gives

$$A_0^2 = \frac{2\hbar\omega}{n_r n_g \varepsilon_0 \omega^2 V}$$
(42)

Substituting Equations, (15), (38), (41), and (42) into Equation (10) gives

$$R_{sp}(\hbar\omega) = \frac{1}{\hbar\omega} \frac{\pi e^2 \hbar}{n_r n_g \varepsilon_0 m_0^2} \frac{2}{3} |M|^2 \rho_{red} (E_{eh} - E'_g) \rho_{opt}(\hbar\omega) f_c (1 - f_v)$$
(43)

in units of transitions per (s cm<sup>3</sup> energy) (Corzine, 1993: 42-3). This is the equation used to calculate the spontaneous emission spectrum.

## **Intraband Relaxation**

The spectrum calculated by Equation (43) will have sharp peaks resulting from the discrete density of states. In reality intraband relaxation will cause the peaks to broaden. This is due to carrier-carrier scattering and carrier-phonon scattering. This process can be described by convolving the spectrum calculated with Equation (43) with a broadening function B:

$$r_{sp}(\hbar\omega) = \int_{E'_s}^{\infty} R_{sp}(E_{eh}) B(\hbar\omega - E_{eh}) dE_{eh}$$
(44)

where  $r_{sp}(\hbar\omega)$  is the convolved spontaneous emission as a function of the photon energy,  $\hbar\omega$  and  $R_{sp}(E_{eh})$  is the unconvolved spontaneous emission as a function of the energy difference between the electron and hole states (Asada, 1993: 99). The broadening function used is derived in Asada, 1993. The lineshape used is non-Lorentzian and comes from the Fourier transform of the response of the polarization to an impulse electric field (Asada, 1993: 98).

It should be noted here that the intraband relaxation convolution is not the convolution mentioned in the Introduction. The intraband relaxation convolution takes into account a phenomenon that is inherent to the semiconductor structure. The convolution mentioned in the Introduction is performed to take into account a phenomenon that is introduced in the measurement process. This phenomenon will be described in the next Chapter. The terms "calculated spectrum" and "calculated widths" used in this thesis refer to spectra calculated with Equation (44) and the width of these spectra. The terms "convolved calculated spectra" and "convolved calculated widths" refer to the spectra that have been corrected for the experimental broadening mentioned in the Introduction and the widths of these spectra.

## **Summary of Calculation Procedure**

The goal of the theoretical calculations is to determine the width of the PL spectra that are expected for different carrier densities. The calculation involves several steps. First the energy levels of the well region are calculated using Equations (5). Then spontaneous emission spectra are calculated for a range of carrier densities using Equation (43). The spectra are then convolved according to Equation (44) to account for intraband relaxation. Plots of some of these spectra will be presented in Chapter 4.

As explained in the next chapter, these spectra have to be convolved again to account for broadening that occurs in the experimental measurements before the widths can be determined. Then the widths of the spectra can be plotted as a function of the carrier density used to calculate each PL spectrum.

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# 3. Experiment

## Overview

Capt Craig Largent of the Department of Engineering Physics at AFIT took the experimental data used in this thesis using the UMIPS constructed by Capt Cooley. This chapter will describe the experiment so the reader will understand what was measured and how the data was taken. The goal of the measurements is to obtain the width of the PL spectra at different delay times. The next section describes the configuration of the UMIPS and the technique. The third section describes the upconversion process and the fourth section describes efficiency and tuning considerations. The fifth section describes the method used to measure the spectra and presents the results as a plot of measured width as a function of delay time. The last section discusses the physical source of the convolution discussed in the Introduction.

## **UMIPS Experiment**

The quantum well sample is optically pumped by a pulse of laser light. Figure 14 shows a schematic of the UMIPS. Using a mode locked laser, short pulses can be produced at high repetition rates (Verdeyen, 1995: 296-7). The pump beam is split so that only one leg reaches the sample. The sample is excited and spontaneous emission occurs. The emission is recombined with the second leg of the pump beam in a Potassium Titanyl Arsenate (KTA) crystal. Sum frequency generation (SFG), or upconversion, occurs producing a third wave at a different wavelength, which is measured by a spectrometer.



Upconversion will take place in the crystal and produce a third beam of

upconverted photons such that

$$\omega_p + \omega_{PL} = \omega_{up} \tag{45}$$



where  $\omega_p$ ,  $\omega_{PL}$ , and  $\omega_{up}$  are the frequencies of the pump beam, the PL wave, and the upconverted wave, respectively. Upconversion will only take place when the pump and PL waves overlap in space and time in the nonlinear optical crystal. As shown in Figure 15, the pump beam acts as a light gate. By varying the distance the upconversion pump beam has to travel, the PL can be sampled at different delay times after the sample has been pumped (Shah, 1988: 277).

## Upconversion

Upconversion is a nonlinear optical phenomena. When an electromagnetic wave interacts with a medium, the polarization induced in the medium can be expressed as a series expansion

$$P_i = \varepsilon_0 \chi_{ij} E_j + 2d_{ijk} E_j E_k + \cdots$$
(46)

where  $P_i$  is the induced polarization vector,  $E_i$  is the induced electric field,  $\varepsilon_0$  is the permeability of free space,  $\chi_{ij}$  is the linear susceptibility, and  $d_{ijk}$  is the second-order susceptibility (Yariv, 1984: 504). Higher order terms in Equation (46) have been dropped.

Upconversion occurs when two waves mix in a crystal for which  $d_{ijk} \neq 0$ . The waves are modeled as plane waves with frequencies  $\omega_1$  and  $\omega_2$ . Plugging these waves into the second term on the right hand side of Equation (46) gives rise to a cross term with a frequency  $\omega_3$ , where  $\omega_3 = \omega_1 + \omega_2$ . The crystal is assumed to be homogeneous, nonabsorbing, and magnetically isotropic (Yariv, 1984: 70). The crystal must be transparent to the three frequencies of interest. The polarization of this new wave is given by

$$P_{i}^{\omega_{3}} = 2d_{ijk}E_{i}^{\omega_{1}}E_{k}^{\omega_{2}}$$
(47)

The exact form of the susceptibility tensor,  $d_{ijk}$ , depends on the crystal symmetry (Yariv, 1984: 506-7).

## Efficiency

The efficiency of the upconversion process affects the strength of the upconverted signal. For the case of negligible pump depletion this is given as

$$\eta(0) = \frac{2 \pi^2 d_{eff}^2 L^2(P_p/A)}{c \varepsilon_0^3 \lambda_{PL} \lambda_{up} n_p n_{PL} n_{up}}$$
(48)

where  $\eta(0)$  is the quantum efficiency,  $d_{eff}$  is the effective nonlinear coefficient of the crystal, L is the length of the interaction in the crystal,  $P_p$  and A are the power and area of the pump beam on the crystal, and  $n_p$ ,  $n_{PL}$ , and  $n_{up}$  are the indexes of refraction for the pump, PL, and upconverted beams respectively (Shah, 1988: 278). The waves must be in

phase with each other or they will interfere and the efficiency will be reduced. The wave vectors are governed by

$$\mathbf{k}_{up} = \mathbf{k}_{p} + \mathbf{k}_{PL} - \Delta \mathbf{k} \tag{49}$$

where phase matching is achieved when  $\Delta \mathbf{k} = 0$  and Equation (45) is valid (Midwinter, 1965: 1135). The quantum efficiency will decrease with increasing  $\Delta \mathbf{k}$  as

$$\eta(\Delta \mathbf{k}) = \eta(0) \frac{\sin^2(L \Delta \mathbf{k})}{(L \Delta \mathbf{k})^2}$$
(50)

where  $\eta(0)$  is the efficiency for the phase matched case and is defined in Equation (48) (Shah, 1988: 278). The phase matching angle can be calculated for this interaction by adding the pump and PL wave vectors according to Equation (49) as shown in Figure 16. Using the law of cosines and  $k = 2\pi n/\lambda$  gives

$$\frac{n_{up}^2}{\lambda_{up}^2} = \frac{n_{PL}^2}{\lambda_{PL}^2} + \frac{n_p^2}{\lambda_p^2} - 2\frac{n_{PL}}{\lambda_{PL}}\frac{n_p}{\lambda_p}\cos(\pi - \psi)$$
(51)

where  $\psi$  is the angle between the pump and PL wave vectors in the crystal in radians (Cooley, 1996: 2-22).

The KTA crystal is biaxial and is oriented so that phase matching will occur when the ordinary polarization of the PL mixes with the extraordinary polarization of the pump beam to produce an upconversion wave that is polarized in the ordinary direction. This is



called an "oeo" interaction and the index of refraction for the pump depends on the angle between the pump beam and the optic axis. Since  $n_p$  is a function of  $\theta$ , this equation can be solved for the angle that the pump beam will make with the optic axis so that  $\Delta \mathbf{k}$  is zero. This angle is the phase matching angle and will change as the wavelengths of the pump and PL change. Thus by turning the crystal the upconversion can be tuned to a specific wavelength. Also, for the oeo interaction  $d_{eff} = d_{32} \sin \theta_{up}$ , where  $\theta_{up}$  is the angle between the upconverted wave vector and the optic axis and  $d_{32}$  comes from the nonlinear susceptibility tensor (Dmitriev, 1991: 24).

#### Measured Spectral Widths

Capt Largent used the UMIPS to measure the upconversion spectra as a function of delay time. Data can be taken by scanning in either wavelength or time. The spectrometer can be scanned over a range of wavelengths for a set delay time. Then the delay time is adjusted and the spectrometer scans over the wavelength range at the new time delay. PL spectra can be measured at each time delay this way. As was mentioned earlier in the Chapter, the crystal is tuned to a wavelength of interest by turning it to a phase matching angle. If the spectrometer is scanned and the crystal is not tuned to the wavelength the spectrometer is set for, the intensity of the signal will decrease for wavelengths that are not near the wavelength the crystal is tuned to. Since the UMIPS does not have a rotation stage that can turn in step with the spectrometer, the spectra were measured by setting the spectrometer to a wavelength of interest, tuning the crystal to that wavelength, and scanning the delay stage over a range of time. By repeating this for several wavelengths at the same delay times, a PL spectrum can be found by taking all of the data for a particular delay time and plotting it as a function of wavelength.

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Data was taken at 77 K by setting the spectrometer to a set wavelength and turning the KTA crystal to achieve the maximum signal. Thirty spectra were measured at delay times ranging from 100 ps to 3 ns in steps of 100 ps. Some of the spectra for different delay times are plotted in Figure 17.

To find the width of the spectra, each point was converted from wavelength to energy space. Capt Largent fit the data in each time slice with a sixth order polynomial and the FWHM of the fit was calculated. The results are shown in Figure 18. As expected the widths decrease with time. This indicates the carrier density is dropping due to recombination as time passes.

## **Broadening Due To Upconversion**

When the pump and PL waves mix in the KTA crystal, no broadening of the upconversion wave will occur if the pump beam is a delta function. By design the pump laser has been set up to emit short pulses to maintain the temporal resolution of the





upconversion spectra. This means that the spectrum in frequency space is large and the upconversion spectrum will be the convolution of the pump and PL spectra. This is because the entire pump spectrum can add with the first energy of the PL spectrum and with the second energy in the pump spectrum and so on. To account for this effect, the calculated spectra were convolved with a measured spectrum of the pump beam so that an accurate comparison of the calculated and measured widths could be performed.

## 4. Results

## **Overview**

This chapter presents the results of the calculations and compares them to previously measured data. First the sample is described and the material parameters for the well and barrier discussed. The third section discusses the calculation of the spontaneous emission spectral widths as a function of total carrier density. The fourth section explains the convolution procedure used to adjust the calculated spectral widths for broadening that occurs in the measurement process. The last section shows the carrier density as a function of time and compares this result to Cooley's result.

### Sample Description

The sample was a multiple quantum well structure consisting of ten InAs<sub>0.935</sub>Sb<sub>0.065</sub> well layers ten nm thick interspersed between 11 In<sub>0.85</sub>Al<sub>0.15</sub>As<sub>0.9</sub>Sb<sub>0.1</sub> barrier layers that are 20 nm thick. The sample was grown by Molecular Beam Epitaxy on an InAs substrate at the Massachusetts Institute of Technology/Lincoln Laboratory (MIT/LL). The sample has been thoroughly described in a previous publication (wafer B in Choi, 1996). Details of the growth process have also been published previously (Turner, 1995). Figure 19 shows an energy diagram of the laser structure. A conduction band offset of 0.75 was used in the calculations. This number came from Cooley who got it from Turner.

Several parameters are needed for the calculations that depend on the type of materials used in the laser structure. The well material is the most important in terms of its effect on the calculations. This sample has a ternary structure of the form  $InAs_{1-x}Sb_x$ 

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Figure 19: Quantum well structure for Laser B (after Figure 4-1 in Cooley, 1996: 4-2).

which has not been thoroughly characterized. Where values for the ternary have not been published, linear interpolation between the values for InAs and InSb was used to find a value for the sample. Table 1 lists the parameters for the binary compounds used and the interpolated values found for the well.

Several sources listed empirical relations or data for the energy gap of InAs<sub>1-x</sub>Sb<sub>x</sub> as a function of antimony mole fraction for a given temperature (Woolley, 1964: 1883; Stringfellow, 1971: 805; Osbourn, 1984: 176; Yen, 1987: 928; Fang, 1990: 7038). A relationship for the energy gap as a function of antimony mole fraction and temperature initially published by Wieder and Clawson (Wieder, 1973: 220) and reprinted in Rogalski, 1989: 37, that fits these empirical relations well is

$$E_g(x,T) = 0.411 - \frac{3.4 \times 10^{-4} T^2}{210 + T} - 0.876x + 0.70x^2 + 3.4 \times 10^{-4} xT(1-x)$$
(52)

Parameter	InAs	InSb	AlAs	AlSb	Well	Barrier
$a_0$ (Å)	6.0583	6.47937			6.0857	
$C_{11} (10^{11} \text{ dynes/cm}^2)$	8.329	6.669			8.2211	
$C_{12} (10^{11} \text{ dynes/cm}^2)$	4.526	3.645			4.469	
<i>a</i> (eV)	-6.0ª	-7.7 <sup>a</sup>			-5.9	
<i>b</i> (eV)	<b>-</b> 1.8 <sup>a</sup>	-2.0 <sup>a</sup>			-1.8	
$m_{ m e} \left( { m m_0}  ight)$	0.0239	0.01359	0.15	0.259	0.0232	0.044
$m_{ m hhz}$ (m <sub>0</sub> )	0.35	0.34	0.409	0.336	0.35	0.36
$m_{ m lhz}({ m m_0})$	0.026	0.0158	0.153	0.123	0.025	0.044
$m_{ m hhx}$ (m <sub>0</sub> )	0.35	0.42			0.35	
$m_{\rm lhx}~({ m m_0})$	0.026	0.0158			0.025	
<i>n</i> <sub>r</sub>	3.714 <sup>b</sup>	4.418 <sup>b</sup>	2.875°	3.182	3.760	3.730
$\varepsilon(0)$	15.15	16.8			15.2	

 

 Table 1: Material parameters for binary compounds and interpolated values for the well and barrier. Values that were not calculated or used are left blank.

All values are from Madelung, 1991, except

a) Blacha, 1984

b) Aspnes, 1983

c) Fern, 1971

where x is the antimony mole fraction, T is in Kelvin, and  $E_g$  is in eV. The spontaneous emission spectrum calculated using an energy gap predicted by this relation showed that the gap was to large for this sample as shown in Figure 20.

Major Michael Marciniak studied several  $InAs_{1-x}Sb_x$  samples with varying x and found smaller energy gaps for his samples (Marciniak, 1995). His samples were bulk samples grown by MIT/LL using Molecular Beam Epitaxy on GaSb substrates. Marciniak fit the temperature dependence of the energy gap for his samples using the Varshni relation (Equation 6). Using his data from the samples with x = 0.059 and x = 0.071, I interpolated point by point to find the temperature dependence for the sample I studied (x = 0.065). The interpolated data was fit to the Varshni relation to find  $E_g(0)$ ,  $\alpha$ , and  $\beta$  for wafer B and resulted in

$$E_g(T) = 0.327 - \frac{3.79 \times 10^{-4} T^2}{346 + T}$$
(53)

for x = 0.065 and  $E_g$  in eV. Figure 20 shows a comparison between a time integrated spectrum of the sample and the spectra calculated with Equations (52) and (53) for T = 77K. Equation (53) was used to calculate the well energy gap since it matched the experimental data from the sample better. Marciniak found inconclusive evidence of phase separation in his samples (Marciniak, 1995: 7-47 - 7-48). Since wafer B was grown by the same researchers using the same technique it may suffer from the same problems as Marciniak's samples.

The split-off energy band separation,  $\Delta_{so}$ , was calculated with

$$\Delta_{xx}(x) = 1.17x^2 - 0.75x + 0.39 \tag{54}$$

for x = 0.065 and  $\Delta_{so}$  in eV (Rogalski, 1989: 37). This relation is used in Equation (39) to calculate  $|M|^2$  for the well material.

Table 2 lists the calculated parameters for the system under strain at T = 77 K. The unstrained well energy gap was found using Equation (53). The strain and energy band shifts were calculated with Equations (7) and (8). The spin-orbit splitting and  $|M|^2$  were calculated as described in the last paragraph.



Table 2: Calculated well parameters. Values are inmeV unless stated otherwise.				
$\varepsilon_{\rm s}$ (percent)	0.450			
Well $E_{g}$	334			
$\Delta E_{ m hh}$	11.7 41.3			
$\Delta E_{ m lh}$				
hh-lh separation	29.6			
$\Delta_{ m so}$	346			
$ M ^2 (2 \text{ eV}/m_0)$	16.9			

The barrier material is a quarternary of the form  $In_{1-y}Al_yAs_{1-x}Sb_x$ . Values for most of the barrier parameters A (except  $\alpha$  and  $\beta$ ) were interpolated with

$$A = (1 - x)A(InSb) + (1 - x)(1 - y)A(InAs) + xyA(AISb) + x(1 - y)A(AIAs)$$
(55)

using the values for InAs, InSb, AlAs, and AlSb, where x is the mole fraction of antimony and y is the mole fraction of aluminum (Madelung, 1991: 156). Table 1 lists the values for the binaries used to interpolate the values for the barrier.

The energy gap for the barrier was calculated at a given temperature by calculating the gap for each of the four binary compounds using Equation (6). The Varshni parameters used are listed in Table 3 along with the resulting energy gaps calculated for T= 77 K. These four energy gaps were then interpolated using Equation (55) to find the barrier band gap listed in the table. The fit parameters  $\alpha$  and  $\beta$  were not calculated for the barrier because these parameters are derived from the fit to the Varshni relation. Their values are not related to the mole fractions and so cannot be interpolated (Marciniak, 1995:6-42 to 6-45 discusses this).

energy gap at $T = 77$ K for each binary compound. These energy gaps were interpolated using Equation (55) to find the barrier gap.					
Varshni Parameter	InAs <sup>a</sup>	InSb <sup>a</sup>	AlAs <sup>b</sup>	AlSb <sup>c</sup>	Barrier
$\alpha$ (10 <sup>-4</sup> eV/K)	2.76	2.7	15.35	4.68	
β(K)	83	106	1018	190	
$E_g(0 \text{ K}) (\text{eV})$	0.415	0.235	3.133	2.384	
<i>E<sub>g</sub></i> (77 K) (meV)	405	226	3125	2374	641

Table 3: Barrier energy gap calculation. The Varshni parameters were used to find the

a) Fang, 1990

b) from fit of data in figure from Monemar, 1973

c) from fit of data in figure from Joullie, 1982

## Spectral Width Calculation

The first step in a calculation of the spectral widths is to calculate the energy levels in the quantum well. The depths of the wells were found using Equations (9) with the values in Table 2 and Table 3 and the conduction band offset listed on page 36. These well depths, listed in Table 4, were used in Equation (5) to calculate the energy levels for the conduction, heavy hole, and light hole bands  $(E_{cn}, E_{hhn})$ , and  $E_{lhn}$ ) numerically. The energy levels are listed in Table 5 and plotted with the wells in Figure 21. The Mathematica code used in this calculation is listed in Appendix A.

Table 4: Well depth	s in meV at $T = 77$ K.	Table 5: Energy levels	in meV at $T = 77$ K.
V <sub>cb</sub>	231	CB Level 1	55.1
$V_{ m hh}$	76.8	CB Level 2	210.7
$V_{ m lh}$	47.2	HH Level 1	7.0
		HH Level 2	27.2
		HH Level 3	58.2
		LH Level 1	54.0



The unconvolved spontaneous emission spectrum is calculated next. The well energies are used to define where the steps in the density of states occur (see Figure 9 on page 17). The quasi-Fermi levels  $E_{Fc}$  and  $E_{Fv}$  were found by solving Equation (31) numerically for the conduction band and the valence band for a given carrier density. The well energies, quasi-Fermi levels, and  $|M|^2$  were then used to calculate the spontaneous emission spectrum for a given temperature and total carrier density with Equation (43). The spectra were convolved using Equation (44), which finishes the computation of the calculated spectra. The MatLab files used to calculate the spectra are listed in Appendix B. They are based on code written by Chia-Fu Hsu, a Ph.D. student of Peter Zory at the University of Florida. Figure 22 shows a plot of some calculated spectra for different carrier densities.

The widths of the spectra were calculated using the Mathematica code in Appendix C. One hundred spectra were calculated with carrier densities ranging from 1 x



Figure 22: Calculated PL spectra for different carrier densities. The carrier densities are in units of cm<sup>-3</sup> and were multiplied by the factors shown to fit them on the plot.

10<sup>16</sup> cm<sup>-3</sup> to 1 x 10<sup>18</sup> cm<sup>-3</sup>. Each calculated spectrum is composed of 625 discrete points. The energy spacing (along the x axis) between these points is 1 meV. Decreasing this spacing increased calculation time dramatically, so the FWHM of the spectra was found by linearly interpolating between the two points nearest to the half maximum value at each side of the peak in order to gain more resolution. Each of these 100 spectra were calculated for a total carrier density, so the FWHM of these emission spectra can be plotted vs. carrier density.

## **Convolution With Upconversion Pump Beam**

As described in Chapter 3, the upconversion process occurs when the second leg of the pump beam mixes with the PL from the sample in the KTA crystal. The resulting upconversion spectrum is broadened due to the convolution of the two electromagnetic waves. In order to compare the widths of the calculated spectra with the widths of the measured spectra, this effect must be taken into account. This is done by convolving the calculated spectra with the measured spectrum of the pump beam. The code used to perform the convolutions is listed in Appendix C.

The first step was to measure the spectrum of the Ti:Sapphire pump beam while it was mode locked. Rick Patton, a technician with the Engineering Physics Department, performed this measurement by routing the beam into an EG&G model 1471 optical multichannel analyzer using an EG&G model 1453 silicon photodiode detector. The spectrum was converted from wavelength to energy and is shown in Figure 23. The FWHM of the peak is 15.39 meV. The spectrum was fit with a 30<sup>th</sup> order polynomial so that the spectrum could be regenerated with the same energy spacing as the calculated PL spectra.

Convolution is represented by  $\otimes$  and  $h = f \otimes g$  is defined as (Jansson, 1984: 6)



$$h(x) = \int_{\infty}^{\infty} f(x-x')g(x')dx' \quad .$$

(56)

It is well known that the product of the Fourier Transform of two functions, f and g, is the Fourier Transform of the convolution of f and g:

$$\Im(f)\Im(g) = \Im(f\otimes g) = \Im(h)$$
<sup>(57)</sup>

where

$$\Im\left(f(\omega)\right) = \frac{1}{\sqrt{2\pi}} \int_{-\infty}^{\infty} f(\omega) e^{i\omega t} d\omega = f(t)$$
(58)

is the Fourier transform of  $f(\omega)$  (Jansson, 1991: 11). The convolution, h, can be found by taking the inverse Fourier transform of the right hand side of Equation (57).

The convolution was performed using two methods. The first method used Fourier transforms with Equation (57). The high energy end of the calculated PL spectra was dropped to make the list of data 512 elements long. The values of the dropped elements were essentially zero and would have no effect on the convolution. The pump spectrum was regenerated with an energy spacing of 1 meV and zeros were added to each end of the list to make it 512 elements long (an equal number of points was required to use Mathematica's "Fourier" function). The transforms of the pump and calculated PL spectra were multiplied together and the inverse transform of the product taken with Mathematica's "InverseFourier" function. The widths were determined as described in the previous section on page 44.

The second method used Equation (56) to compute the convolution. The PL spectra were not truncated and no extra zeros were added to the pump spectrum. The pump spectrum data list was reversed and lined up with the left side of the calculated PL spectrum. Since the energy spacing of the two spectra is the same, the sum of the energy values (x coordinate) of each pair of lined up data points will be the same. This sum of

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the energy values is the energy (x coordinate) for the first point in the convolved spectra. Each intensity data point (y coordinate) from the reversed pump spectrum was multiplied with the intensity data point (y coordinate) from the PL spectrum that was lined up with it. These products were summed and this was the value of the intensity (y coordinate) of the first data point of the convolved spectrum. The reversed pump was moved one element to the right and the process repeated until the end of the PL spectrum was reached. The end result is the convolved spectrum which corresponds to the upconversion spectra. The widths were determined as described in the previous section on page 44.

Both methods of convolution yielded the same widths for the convolved spectra. Since each of the PL spectra were calculated with a specific carrier density, these new convolved calculated widths can be plotted as a function of the carrier density of the PL spectrum used in the convolution. The calculated widths and convolved calculated widths are plotted as a function of carrier density in Figure 24. The effect of the convolution is



greatest when the PL widths are smaller. As the PL widths widen, the pump beam width becomes narrower in comparison and broadens the PL less.

## **Carrier Density as a Function of Time**

Since the convolved calculated widths were determined for specific carrier densities and the measured widths for specific times it should be possible to map the carrier densities to time. Each point in Figure 18 was measured at a specific time. The width measured at that time can be compared with the convolved widths calculated in Figure 24. Each of these convolved widths was calculated with a specific carrier density, so the a plot of carrier density as a function of time can be created. This plot is shown in Figure 25 and the code used to calculate the data is in Appendix D.

Capt Cooley calculated the carrier density as a function of time using a different method (Cooley, 1996: 5-1 to 5-7). He measured the luminescence from the sample at one wavelength for different delay times and then plotted the luminescence as a function



of time. Using the absorption of the material, he calculated the two dimensional carrier density created by different pump beam powers. He measured the luminescence over a range of times for each of these pump power levels and associated the first data point at each power with his calculated value of two dimensional carrier density. This gave him the luminescence as a function of carrier density. He connected the carrier density to the times through the luminescence and converted from a two dimensional carrier density to a three dimensional carrier density using

$$n_{3D} = \frac{n_{2D}}{NL_z}$$
(59)

where N is the number of quantum wells and  $L_z$  is the thickness of a quantum well. This relationship assumes that the carriers are confined to the well regions.

A comparison of Cooley's results and the results from this thesis are shown in Figure 26. The data is close at lower carrier densities but does not agree with Cooley's results at higher carrier densities. This discrepancy could be due to a slight change in the



experimental conditions. For example, the spot size of the pump beam on the sample could have changed slightly between the times the measurements were taken. This will affect the number of carriers created and Cooley's absorption calculation. Also the model I used to calculate the PL spectra assumes parabolic bands. This assumption is good near the band edge and becomes less accurate the farther the carriers are from the edge. At higher carrier densities higher levels of the band will be occupied and the applicability of the parabolic band assumption is reduced.

## 5. Conclusion

## Summary

The objective of this research is to study the temporal behavior of the carrier density in a mid-IR quantum well laser. This will help researchers make more efficient lasers that emit at these wavelengths. PL spectra can be calculated for a given carrier density and measured at different times. The calculated spectra can be compared with the measured spectra to associate a specific carrier density with a specific time. The width of the PL spectra was the parameter chosen to compare spectra.

The quasi-Fermi levels in a semiconductor can be related to the carrier density. As the carrier density increases the PL spectrum will become wider and shift slightly toward higher energies as the higher energy levels become more populated. The energy levels were calculated for an InAs<sub>0.935</sub>Sb<sub>0.065</sub>/In<sub>0.85</sub>Al<sub>0.15</sub>As<sub>0.9</sub>Sb<sub>0.1</sub> multiple quantum well sample at a temperature of 77 K assuming parabolic bands. Using these energy levels 100 spectra were calculated for different carrier densities.

The UMIPS can measure spectra with a time resolution of the width of the pump pulse by measuring the upconversion spectra at different delay times. The PL spectrum of the sample was measured at 30 times using this technique. Because the pulse width of the pump is narrow, its spectrum is wide and the upconversion spectrum will be a convolution of the pump and the PL. The calculated spectra were convolved with the measured upconversion pump beam spectrum. The widths of these convolved calculated spectra were compared to the measured spectral widths to find the carrier density as function of time. These results agree with Cooley's results at low carrier densities and disagree at higher carrier densities. This discrepancy could be due to changing experimental conditions or the break down of the parabolic band model at higher carrier densities.

## **Future Research**

Two things can be done to improve the results of this research. The first would be to calculate the band structure of the material more accurately by using, for example, the  $\mathbf{k} \cdot \mathbf{p}$  method. This would increase the accuracy of the spontaneous emission spectra calculation and provide a better relation between width and carrier density. This would also help determine the carrier densities at which the parabolic band approximation is applicable.

A second improvement would be to slave the rotation stage of the KTA crystal to the spectrometer controller. As the spectrometer is scanned the stage would rotate so the optimum phase matching angle was used for each wavelength. This would allow a complete spectrum over wavelength to be taken at once which would provide more data in each time slice. This would improve the accuracy of the fits and provide a better relation between the measured widths and time. If the improvements are significant it could allow entire spectra to be compared instead of just the widths of the spectra. These improvements would allow better estimates of the carrier density as a function of time to be made.

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# **Appendices**

## **Appendix A: Energy Level Calculation**

#### Last Modified: 3 Oct 97

This file calculates the energy levels in a quantum well. The method follows the MatLab code written by Chia-Fu Hsu, a Ph.D. student of Peter Zory at the University of Florida. This code uses Newton's Method instead of the Bisection Method to numerically solve for the eigenenergies. The equations used are equations 68a and 68b on p. 60 of "Quantum Well Lasers", edited by Peter Zory. This model assumes parabolic bands with a single, finite square well.

The first part of the code is used to define the material parameters for the well and barrier materials. The conduction band offset is used to determine the well depths. The second part calculates the energies for a given well width. The well width is listed two places so make sure the appropriate value is current in the Kernel. The third part displays the results.

```
Off[General::"spell1"]
```

## Parameters

T = 77; (\* temperature of solid in K \*)

Universal constants

 $m0 = 9.10910^{-31}; (* \text{ electron mass in kg *})$   $c = 2.99810^{8}; (* \text{ speed of light in m/s *})$   $h = 6.62610^{-34}; (* \text{ Planck's constant in J \cdot s *})$   $\frac{h}{2\pi};$   $kb = 1.38110^{-23}; (* \text{ Boltzmann's Constant in J/K *})$   $e = 1.60210^{-19}; (* \text{ charge of electron *})$   $\epsilon 0 = 8.85410^{-12}; (* \text{ permitivity of free space *})$ 

Varshni formula and quarternary alloy interpolation formula

$$\varepsilon gT[\alpha_{,}, \beta_{,}, \varepsilon g0_{,}, T_{]} := \varepsilon g0 - \frac{\alpha T^{2}}{T + \beta};$$
  
int[AlAs\_, AlSb\_, InAs\_, InSb\_, y\_, x\_] :=  
 $(1 - x) y InSb + (1 - x) (1 - y) InAs + x y AlSb + x (1 - y) AlAs;$ 

 $\varepsilon$  stands for energy in this notebook.

### Well Parameters

The well width is also listed below in the code to make calculations easier.

 $lz = 100 \, 10^{-10}$ ; (\* well width in m \*)

#### ■ Material: InAs<sub>1-x</sub>Sb<sub>x</sub>

The well in composed of  $InAs_{1-x}Sb_x$  on an InAs substrate. Laser B has an Antimony concentration of 6.5%.

```
x = 0.065; (* Antimony concentration fraction *)
 (* Weider, Clawson heavy hole energy bandgap in eV *)
 (*\epsilon gxT[x_t_] = 0.411 - \frac{3.4 \ 10^{-4} \ t^2}{t + 210} - 0.876 \ x + 0.70 \ x^2 + 3.4 \ 10^{-4} \ t \ x \ (1-x);

εgNoStrain = εgxT[x,T];*)

 (* energy gap from Marciniak's data *)
\varepsilongNoStrain = 0.327479 - \frac{3.7947610^{-4} T^2}{T + 346.193}
 (* lattice parameter in Angstroms *)
a0InAs = 6.0583; a0InSb = 6.47937;
a0 = x a0InSb + (1 - x) a0InAs;
 (* substrate lattice parameter in Angstroms *)
a0s = a0InAs;
(* elastic stiffness coefficients in 10<sup>11</sup> dyne/cm<sup>2</sup> *)
cllInAs = 8.329; cllInSb = 6.669;
cll = x cll InSb + (1 - x) cll InAs;
c12InAs = 4.526; c12InSb = 3.645;
c12 = x c12InSb + (1 - x) c12InAs;
(* hydrostatic deformation potential in eV *)
aInAs = -5.8; aInSb = -7.7; a = x aInSb + (1 - x) aInAs;
(* shear deformation potential in eV *)
bInAs = -1.8; bInSb = -2.0; b = x bInSb + (1 - x) bInAs;
(* strain, positive for compression *)
es = (a0 - a0s) / a0;
(* change in heavy hole energy gap *)
\Delta \varepsilon hh = -2 a \varepsilon s \frac{c11 - c12}{c11} + b \varepsilon s \frac{c11 + c12}{c11};
(* change in light hole energy gap *)
\Delta \varepsilon lh = -2 a \varepsilon s \frac{c11 - c12}{c11} - b \varepsilon s \frac{c11 + 2 c12}{c11};
\varepsilon g = \varepsilon g No Strain + \Delta \varepsilon hh;
(* hh - lh gap *)
s = \Delta \varepsilon lh - \Delta \varepsilon hh;
(* electron effective mass in kg *)
meInAs = 0.0239m0; meInSb = 0.01359m0;
me = x meInSb + (1 - x) meInAs;
(* growth direction heavy hole effective mass in kg *)
mhhzInAs = 0.35m0; mhhzInSb = 0.34m0;
mhhz = x mhhzInSb + (1 - x) mhhzInAs;
(* growth direction light hole effective mass in kg *)
mlhzInAs = 0.026 m0; mlhzInSb = 0.0158 m0;
mlhz = x mlhzInSb + (1 - x) mlhzInAs;
(* in plane direction heavy hole effective mass in kg *)
mhhxInAs = 0.35m0; mhhxInSb = 0.42m0;
mhhx = x mhhxInSb + (1 - x) mhhxInAs;
(* in plane direction light hole effective mass in kg *)
```

```
mlhxInAs = 0.026 m0; mlhxInSb = 0.0158 m0;
mlhx = x mlhxInSb + (1 - x) mlhxInAs;
(* index of refraction *)
nInAs = 3.714; nInSb = 4.418;
n = x nInSb + (1 - x) nInAs;
(* static dielectric coefficient *)
eInAs = 15.15; eInSb = 16.8;
\epsilon = x \epsilon InSb + (1 - x) \epsilon InAs;
(* spin orbit splitting energy in eV *)
\Delta so = 1.17 x^2 - 0.75 x + 0.39;
(* transition matrix elements in eV *)
m2 = \left(\frac{m0}{me} - 1\right) \frac{\varepsilon g + \Delta so}{2 (\varepsilon g + \Delta so 2/3)} m0 \varepsilon g;
Print["For the well:"]
Print[" \varepsilon g (no strain) = ", \varepsilon g No Strain]
Print[" strain corrections: es = ", es]
Print[" a0 for well = ", a0, " Å, a0 for substrate = ", a0s, " Å"]
Print[" \varepsilon g = ", \varepsilon g, " eV, s = ", s, " eV"]
Print[" me = ", me/m0, " m0"]
Print[" mhhz = ", mhhz/m0, " m0, mhhx = ", mhhx/m0, " m0"]
Print[" mlhz = ", mlhz/m0, " m0, mlhx = ", mlhx/m0, " m0"]
Print[" n = ", n, ", \epsilon = ", \epsilon, ", M<sup>2</sup> = ", \frac{2}{m0} m2, " m0/2"]
For the well:
 \epsilon g (no strain) = 0.322162
 strain corrections: \epsilon s = 0.00449738
  a0 for well = 6.08567 Å, a0 for substrate = 6.0583 Å
  \epsilon g = 0.333895 \ eV, \ s = 0.0296039 \ eV
me = 0.0232299 m0
mhhz = 0.34935 mO, mhhx = 0.35455 mO
mlhz = 0.025337 m0, mlhx = 0.025337 m0
```

#### Barrier Parameters

 $lzb = 200 10^{-10}$ ; (\* barrier width in m \*)

n = 3.75976,  $\epsilon = 15.2572$ ,  $M^2 = 16.9087 m0/2$ 

### ■ Material: In<sub>1-v</sub>Al<sub>v</sub>As<sub>1-x</sub>Sb<sub>x</sub>

The concentrations are also listed in the well section to calculate the strain on the well.

xb = 0.1; (\* Antimony concentration fraction \*)
yb = 0.15; (\* Aluminum concentration fraction \*)

```
(* barrier lattice parameter in Angstroms *)
a0AlAs = 5.660; a0AlSb = 6.1355;
a0b = int[a0AlAs, a0AlSb, a0InAs, a0InSb, yb, xb]; (* heavy hole energy bandgap in eV *)
\alphaInAs = 2.7610<sup>-4</sup>; \betaInAs = 83; \epsilong0InAs = 0.415;
\epsilongInAs = \epsilongT[\alphaInAs, \betaInAs, \epsilong0InAs, T];
\alphaInSb = 2.7010<sup>-4</sup>; \betaInSb = 106; \epsilong0InSb = 0.235;
\varepsilongInSb = \varepsilongT[\alphaInSb, \betaInSb, \varepsilong0InSb, T];
\alphaAlAs = 15.347 10<sup>-4</sup>; \betaAlAs = 1018; \epsilongOAlAs = 3.133;
\epsilongAlAs = \epsilongT[\alphaAlAs, \betaAlAs, \epsilongOAlAs, T];
\alphaAlSb = 4.68410<sup>-4</sup>; \betaAlSb = 190; \epsilongOAlSb = 2.384;
\epsilongAlSb = \epsilongT[\alphaAlSb, \betaAlSb, \epsilongOAlSb, T];
egb = int[egAlAs, egAlSb, egInAs, egInSb, yb, xb];
(* electron effective mass in kg *)
meAlSb = 0.259m0; meAlAs = 0.15m0; meInAs = 0.0239m0; meInSb = 0.01359m0;
meb = int[meAlAs, meAlSb, meInAs, meInSb, xb, yb];
(* heavy hole effective mass in kg *)
mhhzAlSb = 0.336m0; mhhzAlAs = 0.409m0; mhhzInAs = 0.35m0; mhhzInSb = 0.34m0;
mhhb = int[mhhzAlAs, mhhzAlSb, mhhzInAs, mhhzInSb, xb, yb];
(* light hole effective mass in kg *)
mlhzAlSb = 0.123m0; mlhzAlAs = 0.153m0; mlhzInAs = 0.026m0; mlhzInSb = 0.0158m0;
mlhb = int[mlhzAlAs, mlhzAlSb, mlhzInAs, mlhzInSb, xb, yb];
Print["For the barrier:"]
Print[" a0 = ", a0b, " A"]
Print[" \varepsilon g = ", \varepsilon g b, " eV"]
Print[" me = ", meb/m0, " m0, mhh = ", mhhb/m0, " m0, mlh = ", mlhb/m0, " m0"]
For the barrier:
 a0 = 6.08245 Å
\epsilon g = 0.641398 \, eV
me = 0.0435737 m0, mhh = 0.356905 m0, mlh = 0.043733 m0
```

#### Depths of conduction band and valence band wells

```
\varepsilon \text{coffset} = 0.75; (* \text{ conduction band offset for materials })
\varepsilon \text{ch} = \varepsilon \text{coffset} (\varepsilon \text{gb} - \varepsilon \text{g}); (* \text{ conduction band well depth in eV })
\varepsilon \text{vh} = (1 - \varepsilon \text{coffset}) (\varepsilon \text{gb} - \varepsilon \text{g}); (* \text{ valence band well height in eV })
\text{Print["conduction band well depth} = ", \varepsilon \text{ch}, " eV"]
\text{Print["valence band well height} = 0.230627 \text{ eV}
\text{valence band well height} = 0.0768755 \text{ eV}
```

### List Parameters

```
Print["temperature = ", T, " K"]
 Print["For the well:"]
 Print[" \varepsilong (no strain) = ", \varepsilongNoStrain]
 Print[" strain corrections: es = ", es]
 Print[" a0 for well = ", a0, " Å, a0 for substrate = ", a0s, " Å"]
 Print[" a0 for barrier = ", a0b, " A"]
 Print[" ɛg = ", ɛg, " eV, s = ", s, " eV"]
 Print[" me = ", me/m0, " m0"]
Print[" mhhz = ", mhhz/m0, " m0, mhhx = ", mhhx/m0, " m0"]
Print[" mlhz = ", mlhz/m0, " m0, mlhx = ", mlhx/m0, " m0"]
Print[" n = ", n, ", \epsilon = ", \epsilon, ", M<sup>2</sup> = ", \frac{2}{m0} m2, " m0/2"]
Print["For the barrier:"]
Print[" \varepsilon g = ", \varepsilon g b, " eV"]
Print[" me = ", meb/m0, " m0, mhh = ", mhhb/m0, " m0, mlh = ", mlhb/m0, " m0"]
Print["conduction band well depth = ", \epsilonch, " eV"]
Print["valence band well height = ", ɛvh, " eV"]
temperature = 77 \text{ K}
For the well:
 \epsilon g (no strain) = 0.322162
 strain corrections: \epsilon s = 0.00449738
  a0 for well = 6.08567 Å, a0 for substrate = 6.0583 Å
  a0 for barrier = 6.08245 Å
  \epsilon q = 0.333895 \text{ eV}, \text{ s} = 0.0296039 \text{ eV}
 me = 0.0232299 m0
 mhhz = 0.34935 mO, mhhx = 0.35455 mO
 mlhz = 0.025337 m0, mlhx = 0.025337 m0
 n = 3.75976, \epsilon = 15.2572, M^2 = 16.9087 \text{ mO}/2
For the barrier:
 \epsilon g = 0.641398 \ eV
me = 0.0435737 m0, mhh = 0.356905 m0, mlh = 0.043733 m0
conduction band well depth = 0.230627 eV
valence band well height = 0.0768755 eV
```

# Energy Levels of a single quantum well

For ease of recomputing the well width is listed here so it can be changed.

## $lz = 100 \, 10^{-10}$ ; (\* well width in m \*)

Set up equations 68a and 68b for electrons, heavy holes, and light holes. The equations come from solving Schrödinger's equation and applying boundary conditions. The energy,  $\varepsilon$ , is in meV. The strain separation, s, for the light hole solutions is corrected for in the kzlh expression. The energies used by the light hole equations are really  $\varepsilon$ +s from s to the valence band height. This accounts for the compressive strain that pushes the light holes away from the gap.

Clear[
$$\varepsilon$$
]  
kze =  $\sqrt{\frac{2 \text{ me}}{\hbar^2}} \frac{\varepsilon}{1000} \text{ e}$ ;  $\alpha ze = \sqrt{\frac{2 \text{ meb}}{\hbar^2}} \left(\varepsilon ch - \frac{\varepsilon}{1000}\right) \text{ e}$ ;  
evene = Tan[kze  $\frac{1z}{2}$ ] -  $\frac{\text{me} \alpha ze}{\text{meb kze}}$ ; odde = Cot[kze  $\frac{1z}{2}$ ] +  $\frac{\text{me} \alpha ze}{\text{meb kze}}$ ;  
kzhh =  $\sqrt{\frac{2 \text{ mhz}}{\hbar^2}} \frac{\varepsilon}{1000} \text{ e}$ ;  $\alpha zhh = \sqrt{\frac{2 \text{ mhb}}{\hbar^2}} \left(\varepsilon vh - \frac{\varepsilon}{1000}\right) \text{ e}$ ;  
evenhh = Tan[kzhh  $\frac{1z}{2}$ ] -  $\frac{\text{mhz} \alpha zhh}{\text{mhb} kzhh}$ ; oddh = Cot[kzhh  $\frac{1z}{2}$ ] +  $\frac{\text{mhz} \alpha zhh}{\text{mhb} kzhh}$ ;  
kzlh =  $\sqrt{\frac{2 \text{ mlz}}{\hbar^2}} \left(\frac{\varepsilon}{1000} - s\right) \text{ e}$ ;  $\alpha zlh = \sqrt{\frac{2 \text{ mlb}}{\hbar^2}} \left(\varepsilon vh - \frac{\varepsilon}{1000}\right) \text{ e}$ ;  
evenlh = Tan[kzlh  $\frac{1z}{2}$ ] -  $\frac{\text{mlz} \alpha zlh}{\text{mlbb} kzlh}$ ; oddlh = Cot[kzlh  $\frac{1z}{2}$ ] +  $\frac{\text{mlz} \alpha zlh}{\text{mlbb} kzlh}$ ;

Plot the even and odd solutions to find guesses for the root finding. If the plots have too few points to make a good guess, increment the appropriate PlotData table to calculate more points. This is especially helpful for the heavy hole plots. It is easier to see if a plot crosses the axis if the plots are temporarily made larger by draging the frame. The plotrange can also be varied so the plots will show more points in the lines. The xxPlotData tables are used to plot the functions with discrete points. This avoids having intersections with the energy axis at the assymptotes of the trigonometric functions. The lower limit for the light holes is s meV + 1 meV to prevent kzlh =  $\sqrt{0}$ . This occurs when  $\varepsilon = s$ .

```
plotrange = 5;
 Clear[evenePlotData, oddePlotData,
  evenhhPlotData, oddhhPlotData, evenlhPlotData, oddlhPlotData]
 evenePlotData = Table[{ɛ, evene}, {ɛ, 1, 1000 ɛch}];
 oddePlotData = Table[{ɛ, odde}, {ɛ, 1, 1000 ɛch}];
 evenhhPlotData = Table[{\varepsilon, evenhh}, {\varepsilon, 1, 1000 \varepsilonvh, 0.5}];
oddhhPlotData = Table[{ɛ, oddhh}, {ɛ, 1, 1000 ɛvh}];
evenlhPlotData = Table [{\varepsilon, evenlh}, {\varepsilon, 1000 s + 1, 1000 \varepsilonvh}];
oddlhPlotData = Table[{\varepsilon, oddlh}, {\varepsilon, 1000 s + 1, 1000 \varepsilonvh}];
oddePlot = ListPlot[oddePlotData,
   PlotRange → {-plotrange, plotrange}, AxesLabel → {"meV", "Odd"},
  PlotLabel → FontForm["Electrons", {"Arial", 12}], DisplayFunction → Identity];
evenePlot = ListPlot[evenePlotData,
  PlotRange \rightarrow {-plotrange, plotrange}, AxesLabel \rightarrow {"meV", "Even"},
  PlotLabel → FontForm["Electrons", {"Arial", 12}], DisplayFunction → Identity];
oddhhPlot = ListPlot[oddhhPlotData,
  PlotRange → {-plotrange, plotrange}, AxesLabel → {"meV", "Odd"},
  PlotLabel → FontForm["Heavy Hole", {"Arial", 12}], DisplayFunction → Identity];
evenhhPlot = ListPlot[evenhhPlotData,
  PlotRange → {-plotrange, plotrange}, AxesLabel → {"meV", "Even"},
  PlotLabel → FontForm["Heavy Hole", {"Arial", 12}], DisplayFunction → Identity];
oddlhPlot = ListPlot[oddlhPlotData,
  PlotRange → {-plotrange, plotrange}, AxesLabel → {"meV", "Odd"},
  PlotLabel \rightarrow FontForm["Light Hole", {"Arial", 12}], DisplayFunction \rightarrow Identity];
evenlhPlot = ListPlot[evenlhPlotData,
  PlotRange → {-plotrange, plotrange}, AxesLabel → {"meV", "Even"},
  PlotLabel \rightarrow FontForm["Light Hole", {"Arial", 12}], DisplayFunction \rightarrow Identity];
Print["Well Width = ", lz/10<sup>-10</sup>, " Å, T = ", T, " K"]
Show[GraphicsArray]
  {{oddePlot, evenePlot}, {oddhhPlot, evenhhPlot}, {oddlhPlot, evenlhPlot}}],
DisplayFunction \rightarrow $DisplayFunction]
```

Well Width = 100 Å, T = 77 K


- GraphicsArray -

The equations are solved when  $\varepsilon$  equals the values of the x-intercepts. Plug the guesses from the plots into the respective lists below. Surround the values with curly brackets { } and separate multiple guesses with a comma. If there is no solution set the value equal to {}.

```
oddeGuess = {210}; eveneGuess = {60}; oddhhGuess = {27}; evenhhGuess = {7, 60};
oddlhGuess = {}; evenlhGuess = {57};
```

The eigenvalues are found using Newton's Method for each guess.

```
maxOddE = Length[oddeGuess]; maxEvenE = Length[eveneGuess];
maxOddHH = Length[oddhhGuess]; maxEvenHH = Length[evenhhGuess];
maxOddLH = Length[oddlhGuess]; maxEvenLH = Length[evenlhGuess];
 odde = \{\}; evene = \{\}; oddhe = \{\}; evenhe = \{\}; oddhe = \{\}; evenhe =
 Do[solution = FindRoot[odde == 0, {$\varepsilon$, oddeGuess[[i]]}];
   \varepsilon = \varepsilon /. solution; AppendTo[odde\varepsilon, \varepsilon], {i, 1, maxOddE}];
Do[solution = FindRoot[evene == 0, {$\varepsilon$, eveneGuess[[i]]}];
   \varepsilon = \varepsilon /. solution; AppendTo[evene\varepsilon, \varepsilon], {i, 1, maxEvenE}];
Do[solution = FindRoot[oddhh == 0, {$\varepsilon$, oddhhGuess[[i]]}];
   \varepsilon = \varepsilon /. solution; AppendTo[oddhh\varepsilon, \varepsilon], {i, 1, maxOddHH}];
Do[solution = FindRoot[evenhh == 0, {$$, evenhhGuess[[i]]}];
   \varepsilon = \varepsilon /. solution; AppendTo[evenhh\varepsilon, \varepsilon], {i, 1, maxEvenHH}];
Do[solution = FindRoot[oddlh == 0, {$\varepsilon$, oddlhGuess[[i]]}];
   \varepsilon = \varepsilon /. solution; AppendTo[oddlh\varepsilon, \varepsilon], {i, 1, maxOddLH}];
Do[solution = FindRoot[evenlh == 0, {$\epsilon$, evenlhGuess[[i]]}];
   ε = ε /. solution; AppendTo[evenlhε, ε], {i, 1, maxEvenLH}];
eɛ = Sort[Join[oddeɛ, eveneɛ]];
hhɛ = Sort[Join[oddhhɛ, evenhhɛ]]; lhɛ = Sort[Join[oddhɛ, evenhbɛ]];
Print["Electron Energies: ", eɛ]
Print["Heavy Hole Energies: ", hhɛ]
Print["Light Hole Energies: ", lhɛ]
Electron Energies: {55.0927, 210.669}
Heavy Hole Energies: {6.95165, 27.2429, 58.2015}
Light Hole Energies: {53.9502}
```

## Display Results

This displays the results.

Print["Well Width = ", lz/10<sup>-10</sup>, " Å, T = ", T, " K"] Show[GraphicsArray[ {{oddePlot, evenePlot}, {oddhhPlot, evenhhPlot}, {oddlhPlot, evenlhPlot}}], DisplayFunction → \$DisplayFunction]; Print["Electron Energy Guesses: ", Sort[Join[oddeGuess, eveneGuess]]] Print["Heavy Hole Energy Guesses: ", Sort[Join[oddhhGuess, evenhhGuess]]] Print["Light Hole Energy Guesses: ", Sort[Join[oddlhGuess, evenlhGuess]]] Print["Electron Energies: ", eɛ] Print["Heavy Hole Energies: ", hhɛ] Print["Light Hole Energies: ", lhɛ]

Well Width = 100 Å, T = 77 K



Electron Energy Guesses: {60, 210} Heavy Hole Energy Guesses: {7, 27, 60} Light Hole Energy Guesses: {57} Electron Energies: {55.0927, 210.669} Heavy Hole Energies: {6.95165, 27.2429, 58.2015} Light Hole Energies: {53.9502}

### Well Plot

#### Energy Levels

This plots a diagram of the well and the energy levels.

```
Needs["Graphics`MultipleListPlot`"]
cBand1 = {\{-100, 1000 (\epsilon ch + \epsilon g)\}, \{-49.9, 1000 (\epsilon ch + \epsilon g)\};
cBand2 = Table[{-50+100i, 1000 ɛg}, {i, 0, 1}];
cBand3 = Table[{50.1+49.9i, 1000 (ɛch + ɛɡ)}, {i, 0, 1}];
conductionBand = Join[cBand1, cBand2, cBand3];
vBand1 = Table[{-100+49.9i, -1000 \vert vh}, {i, 0, 1}];
vBand2 = Table[{-50+100i, 0}, {i, 0, 1}];
vBand3 = Table[{50.1+49.9i, -1000 evh}, {i, 0, 1}];
valenceBand = Join[vBand1, vBand2, vBand3];
cbLevel1 = Table[{-50+100i, ec[[1]]+1000cg}, {i, 0, 1}];
cbLevel2 = Table[{-50+100i, eɛ[[2]]+1000ɛg}, {i, 0, 1}];
hhLevel1 = Table[{-50+100i, -hhc[[1]]}, {i, 0, 1}];
hhLevel2 = Table[{-50+100i, -hhɛ[[2]]}, {i, 0, 1}];
hhLevel3 = Table[{-50+100i, -hhc[[3]]}, {i, 0, 1}];
lhLevel1 = Table[{-50+100i, -lhɛ[[1]]}, {i, 0, 1}];
Print["temperature = ", T, " K"]
MultipleListPlot[
 conductionBand, valenceBand, cbLevel1, cbLevel2, hhLevel1, hhLevel2, hhLevel3,
 lhLevel1, PlotJoined → True, PlotStyle → {AbsoluteThickness[2], AbsoluteThickness[2],
   AbsoluteThickness[1], AbsoluteThickness[1], AbsoluteThickness[1],
   AbsoluteThickness[1], AbsoluteThickness[1], AbsoluteDashing[{2, 2}]},
 SymbolShape \rightarrow None, AspectRatio \rightarrow 1.2,
Axes → False, Frame → True, FrameLabel → {"A", "meV", "", ""}];
Print["well \varepsilon g = ", \varepsilon g, " eV, barrier \varepsilon g = ", \varepsilon gb, " eV"]
Print["conduction band well depth = ", \varepsilonch, " eV"]
Print["valence band well height = ", \varepsilon vh, " eV"]
Print["Electron Energies: ", eɛ, " meV"]
Print["Heavy Hole Energies: ", hhɛ, " meV"]
Print["Light Hole Energies: ", lhɛ, " meV"]
temperature = 77 \text{ K}
```



#### Wavefunctions

This calculates the k=0 wavefunctions for the single quantum well. Define a function for k.

$$k[m_{, \varepsilon_{, v_{}}] := \sqrt{\frac{2 m}{\hbar^2} \frac{(\varepsilon - v)}{1000} e}$$

Calculate wavefunctions and load them into tables. The factor of 50 enlarges the scale of the wavefunctions.

cbi¥1 = Table[{d = -100 + i, 50 
$$\frac{\cos[k[me, ec[(1]], 0] (-5010^{-10}]]}{2}$$
  
 $\frac{Exp[-k[meb, ec[(1]], 1000 cch] d10^{-16}]}{2}$  + 1000 cg + cc[[1]]}, (i, 0, 50, 1)];  
cbl¥2 =  
Table[{d = -50 + i, 50  $\frac{\cos[k[me, ec[(1]], 0] d10^{-10}]}{2}$  + 1000 cg + cc[[1]]}, (i, 0, 100, 1)];  
cbl¥3 = Table[{d = 50 + i, 50  $\frac{\cos[k[me, ec[(1]], 0] d10^{-10}]}{2}$  + 1000 cg + cc[[1]]}, (i, 0, 50, 1)];  
Exp[ik[meb, ec[(1]], 1000 cch] d10^{-10}]}{2} + 1000 cg + cc[[1]]}, (i, 0, 50, 1)];  
cbl¥1 = Table[{d = -100 + i, 50  $\frac{\sin[k[me, ec[(2]], 0] (-5010^{-10})]}{2}$   
 $\frac{Exp[ik[meb, ec[(2]], 1000 cch] d10^{-10}]}{2}$  + 1000 cg + cc[[2]]}, (i, 0, 50, 1)];  
cbl¥2 =  
Table[{d = -100 + i, 50  $\frac{\sin[k[me, ec[(2]], 0] d10^{-10}]}{2}$  + 1000 cg + cc[[2]]}, (i, 0, 50, 1)];  
cbl¥2 =  
Table[{d = -50 + i, 50  $\frac{\sin[k[me, ec[(2]], 0] d10^{-10}]}{2}$  + 1000 cg + cc[[2]]}, (i, 0, 50, 1)];  
cbl¥3 = Table[{d = -50 + i, 50  $\frac{\sin[k[me, ec[(2]], 0] d10^{-10}]}{2}$  + 1000 cg + cc[[2]]}, (i, 0, 50, 1)];  
cbl¥3 = Table[{d = -50 + i, 50  $\frac{\cos[k[mha, hac[(1]], 0] d10^{-10}]}{2}$  + 1000 cg + cc[[2]]}, (i, 0, 50, 1)];  
Exp[ik[meb, ec[[2]], 1000 cch] d10^{-10}]} + 1000 cg + cc[[2]]}, (i, 0, 50, 1)];  
Exp[ik[meb, ec[[2]], 1000 cch] d10^{-10}]} - hhc[[1]], (i, 0, 50, 1)];  
Exp[ik[meb, ec[[2]], 1000 cch] d10^{-10}]} - hhc[[1]], (i, 0, 50, 1)];  
hhl¥1 = Table[{d = -100 + i, 50  $\frac{\cos[k[mha, hhc[[1]], 0] d10^{-10}]}{2} - hhc[[1]], (i, 0, 50, 1)];$   
hhl¥2 =  
Table[{d = -50 + i, 50  $\frac{\cos[k[mha, hhc[[1]], 0] d10^{-10}]}{2} - hhc[[1]], (i, 0, 50, 1)];$   
Exp[ik[mhb, hhc[[1]], 1000 evh] d10^{-10}]  
Exp[ik[mhb, hhc[[1]], 1000 evh] d10^{-10}]  
Exp[ik[mhb, hhc[[2]], 1000 evh] d10^{-10}]  
Exp[ik[mhb,

$$\frac{\exp[-i k[mhhb, hhc[[3]], 1000 evh] d 10^{-10}]}{\exp[-i k[mhhb, hhc[[3]], 1000 evh] (-50 10^{-10})]} - hhc[[3]]}, \{i, 0, 50, 1\}];$$
hh3\psi2 =
Table[{d = -50 + i, 50  $\frac{\cos[k[mhhz, hhc[[3]], 0] d 10^{-10}]}{2} - hhc[[3]]}, \{i, 0, 100, 1\}];$ 
hh3\psi3 = Table[{d = 50 + i, 50  $\frac{\cos[k[mhhz, hhc[[3]], 0] 50 10^{-10}]}{2}$ 

$$\frac{\exp[i k[mhhb, hhc[[3]], 1000 evh] d 10^{-10}]}{\exp[i k[mhhb, hhc[[3]], 1000 evh] d 10^{-10}]} - hhc[[3]]}, \{i, 0, 50, 1\}];$$
lh1\psi3 = Table[{d = -100 + i, 50  $\frac{\cos[k[mhhz, hhc[[1]], 0] (-50 10^{-10})]}{2}$ 

$$\frac{\exp[-i k[mhhb, hhc[[1]], 1000 evh] d 10^{-10}]}{2} - hc[[1]]}, \{i, 0, 50, 1\}];$$
lh1\psi2 =
Table[{d = -50 + i, 50  $\frac{\cos[k[mhhz, hhc[[1]], 0] d 10^{-10}]}{2} - hc[[1]]}, \{i, 0, 50, 1\}];$ 
lh1\psi3 = Table[{d = 50 + i, 50  $\frac{\cos[k[mhhz, hhc[[1]], 0] d 10^{-10}]}{2} - hc[[1]]}, \{i, 0, 50, 1\}];$ 
lh1\psi3 = Table[{d = 50 + i, 50  $\frac{\cos[k[mhhz, hhc[[1]], 0] d 10^{-10}]}{2} - hc[[1]]}, \{i, 0, 100, 1\}];$ 
lh1\psi3 = Table[{d = 50 + i, 50  $\frac{\cos[k[mhhz, hhc[[1]], 0] d 10^{-10}]}{2} - hc[[1]]], \{i, 0, 50, 1\}];$ 
lh1\psi3 = Table[{d = 50 + i, 50  $\frac{\cos[k[mhhz, hhc[[1]], 0] d 10^{-10}]}{2} - hc[[1]]], \{i, 0, 50, 1\}];$ 
lh1\psi3 = Table[{d = 50 + i, 50  $\frac{\cos[k[mhhz, hhc[[1]], 0] d 10^{-10}]}{2} - hc[[1]]], \{i, 0, 50, 1\}];$ 
lh1\psi3 = Table[{d = 50 + i, 50  $\frac{\cos[k[mhhz, hhc[[1]], 0] d 10^{-10}]}{2} - hc[[1]]], \{i, 0, 50, 1\}];$ 
lh1\psi3 = Table[{d = 50 + i, 50  $\frac{\cos[k[mhhz, hhc[[1]], 0] d 10^{-10}]}{2} - hc[[1]]], \{i, 0, 50, 1\}];$ 
lh1\psi2 =
Table[i k[mhhb, hhc[[1]], 1000 evh] d 10^{-10}] + hc[[1]]], [i, 0, 50, 1];
htp: join[cbl\psi, cbl\psi2, cbl\psi3]; cbl\psi = Join[chl\psi], cbl\psi2, cbl\psi]; h1\psi2 = Join[cbl\psi], cbl\psi2, cbl\psi3]; h1\psi2 = Join[hh\psi], h1\psi3]; h1\psi3];

Plot wavefunctions.

```
Print["temperature = ", T, " K"]
MultipleListPlot[conductionBand, valenceBand, cbLevel1, cbLevel2, hhLevel1, hhLevel2,
hhLevel3, lhLevel1, cbl\psi, cb2\psi, hhl\psi, hh2\psi, hh3\psi, lh1\psi, PlotJoined \rightarrow True,
 PlotStyle → {AbsoluteThickness[2], AbsoluteThickness[2], AbsoluteThickness[1],
   AbsoluteThickness[1], AbsoluteThickness[1], AbsoluteThickness[1],
   AbsoluteThickness[1], AbsoluteDashing[{2, 2}], AbsoluteThickness[1],
   AbsoluteThickness[1], AbsoluteThickness[1], AbsoluteThickness[1],
   AbsoluteThickness[1], AbsoluteDashing[{2, 2}]}, SymbolShape → None,
AspectRatio → 1.2, Axes → False, Frame → True, FrameLabel → {"A", "meV", "", ""}]
Print["well \varepsilon g = ", \varepsilon g, " eV, barrier \varepsilon g = ", \varepsilon gb, " eV"]
Print["conduction band well depth = ", ɛch, " eV"]
Print["valence band well height = ", $\varepsilon vh, " eV"]
Print["Electron Energies: ", eɛ, " meV"]
Print["Heavy Hole Energies: ", hhɛ, " meV"]
Print["Light Hole Energies: ", lhɛ, " meV"]
```

```
temperature = 77 \text{ K}
```



- Graphics -

well ɛg = 0.333895 eV, barrier ɛg = 0.641398 eV conduction band well depth = 0.230627 eV valence band well height = 0.0768755 eV Electron Energies: {55.0927, 210.669} meV Heavy Hole Energies: {6.95165, 27.2429, 58.2015} meV Light Hole Energies: {53.9502} meV

#### **Appendix B: Spontaneous Emission Spectra Calculation**

Sponlasb.m is the main program used to calculate the spectra. It calls P\_InAsSb.m and P\_BarB.m to input the material parameters for the well and the barrier. It also calls efc.m and efv.m to help with the quasi-Fermi level calculations.

SponlasB.m

e0=8.854\*10<sup>(-12)</sup>;

```
%This program is to calculate gain and spontaneous emission
%spectrum for a QW laser structure.
%
% assumptions:(1) parabolic band model (finite well)
          (2) large strain regime in-plane effective mass
%
 (ansiotropic mass) (???)
%
               (3) non-Lorentzian line-shape function (Ch2,
OWL)
8
°
     written by by C.F. Hsu
÷
                                08/05/96
     modified for Laser B
÷
                                08/01/97 by Anthony Franz
     last modified
                           08/26/97
%
%
          specify well/barrier materials for p <material>.m
%
     1.
÷
         specify mole fractions of the materials
     2.
°
     3. specify Gamma, alpha
     4. choose QB=1 for QW, QB=2 for bulk
°
°
     5. temperature T in K
     6. conduction band offset ratio, Ec off, in percent
%
%
     7. Run Mathematica code to find energy levels
         specify Ni(initial carrier density) & Nf(final
°
     9.
carrier density)
%
     10. specify tau(scattering time)
clear
% universal constant
c = 2.998 \times 10^{8};
h = 6.626 \times 10^{(-34)};
h = h/2/pi;
k = 1.381 \times 10^{(-23)};
m0=9.109*10^{(-31)};
e = 1.602 \times 10^{(-19)};
```

```
% permittivity
```

% condition parameters

T=input('Enter temperature (K) --- '); % enter temperature in K % Sb concentration in well x=0.065; % Sb concentration in barrier xb=0.1; yb=0.15; % Al concentration in barrier material parameter (call function pr % .m) mp=p inassb(T,x); % active layer region % Energy band gap (eV) Eg a=mp(1);% refractive index Nr a=mp(2);r1 = mp(3);% gammal  $r2 \ a = mp(4);$ % gamma2 r3 a = mp(5);% gamma3 Me a=mp(6);% electron effective mass (m0) % z-direction HH effective mass (m0) Mhz a=mp(7);Mlz a=mp(8);% z-direction LH effective mass (m0) Mhx a=mp(9);% in-plane HH effective mass (m0) % in-plane LH effective mass (m0) Mlx a = mp(10);S = mp(11);% splitting energy Eg(LH) -Eg(HH) (eV) % transition coefficient M<sup>2</sup> M = 2 = mp(12);% static dielectric constant (e0) ee=mp(13);% barrier layer region mp=p barB(T,xb,yb); Eq b=mp(1);% Energy band gap (eV) Nr b=mp(2);% refractive index r1 b=mp(3);% gamal  $r2 \ b=mp(4);$ % qama2 r3 b=mp(5);% gama3 % effective mass of e Me b=mp(6);% of hh Mh b=mp(7); Ml b=mp(8);% of lh

% device parameters, kept from Jeff Hsu's code

MT=M 2;% basis function momentum matrix element M<sup>2</sup> Gama=0.01; % confinement factor, from Choi et al, Apl.Phys.Lett.68,2936,(1996) alfa=1.7; % internal loss 1/cm eta\_i=0.67; % injection efficiency % potential profile, offset from Bill's thesis Ec off=0.75; % conduction band offset dEc=Ec off\*(Eg b-Eg a);  $dEv=(1-Ec_off)*(Eg b-Eg a);$ % variable limits hv\_i=0.001; % initial Eeh-Eg hv\_f=0.8; % final Eeh-Eg hv\_step=0.001; % step in photon energy dhv=hv i:hv step:hv f; % photon energy vector dhv1=-1\*hv f:hv step:2\*hv f; % photon energy vector % Data files generated at the very end of the code are named % after the carrier density so update these as well \*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\* Ni=201\*10<sup>15</sup>; % initial carrier density (cm<sup>-3</sup>) Nf=250\*10<sup>15</sup>; % final carier density (cm<sup>-3</sup>) N step=1\*10<sup>15</sup>; % step (cm<sup>-3</sup>) \*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\* % Entering quantum well width in A for Laser B da=100; % QW thickness in A % Entering conduction band and valence band for eigenenergies at kt=0 Ecm=input('Enter C subband energies (meV) --- ')/1000; Ehm=input('Enter HH subband energies (meV) --- ')/1000; Elm=input('Enter LH subband energies (mev) --- ')/1000;

```
N C=length(Ecm); % N_C: # of conduction subbands
N_HH=length(Ehm); % N_HH: # of HH subbands
N_LH=length(Elm); % N_LH: # of LH subbands
Eh1=Ehm(1);
in_ch=min(N_C,N_HH); % # of C-HH transitions, k selection
rule
in cl=min(N C,N LH); % # of C-LH transitions, k selection
rule
% material parameter
               % QW thickness
dz=da;
             % electron effective mass
% z-direction HH effective mass
Me=Me a;
Mhz=Mhz a;
              % z-direction LH effective mass
Mlz=Mlz a;
Mhx=Mhx_a; % in-plane HH effective mass
Mlx=Mlx_a; % in-plane LH effective mass
Mrh=Me*Mhx/(Me+Mhx);% in-plane reduced C-HH effective mass
Mrl=Me*Mlx/(Me+Mlx);% in-plane reduced C-LH effective mass
Mrhz=Me*Mhz/(Me+Mhz)% z-direction reduced C-HH effective
mass
Mrlz=Me*Mlz/(Me+Mlz);% z-direction reduced C-LH effective
mass
            % refractive index
nr=Nr a;
Ro=(nr-1)<sup>2</sup>/(nr+1)<sup>2</sup>;% facet reflectivity
           % facet reflectivity
Rb=Ro:
ee=ee*e0;
               % static dielectric constant
eel=nr^2*e0;
               % optical dielectric constant
Eg=Eg_a;
% initial conditions
iN=0;
% start of N loops
for N=Ni:N step:Nf
  iN=iN+1
```

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```
Eyc=0.15;% initial guess of electron fermi level Eyc=Efc-
Ec
  Eyv=-0.01;% initial guess of hole fermi level Eyv=Ev-Efv
  P=N;
  Nm(iN) = N;
% find fermi level Fc,Fv
                 % differential Energy (eV)
  dE=0.00001;
  tol=0.00001;
                  % tolerance
% using Newton method to find the solution of fermi level
                    % assumed former initial guess
  Fcc=1;
  while abs(Eyc-Fcc) > tol
    % if the difference between two consequent initial
quesses is greater than tolerance
                                    % initial guess as input
    yfc=efc(N,Eyc,Me,T,dz,Ecm);
    yfc = (efc(N,Eyc+dE,Me,T,dz,Ecm)-yfc)/dE; % slope
                                   % initial guess becomes
    Fcc=Eyc;
former inital guess
                                        % new initial guess
    Eyc=Eyc-yfc/yfc ;
  end
                                   % Efc-Ec
  Fcm(iN) = Fcc;
               % assumed former initial guess
  Fvv=1;
  while abs(Eyv-Fvv) > tol
    yfc=efv(P,Eyv,Mhx,Mlx,T,dz,Ehm,Elm);
                                                 % N in cm
    yfc = (efv(N,Eyv+dE,Mhx,Mlx,T,dz,Ehm,Elm)-yfc)/dE;
    Fvv=Eyv;
    Eyv=Eyv-yfc/yfc ;
  end
                                   % Ev-Efv
  Fvm(iN) = Fvv;
  Phm=Mhx*m0*k*T/(pi*h ^2*dz*10^(-10))*log(1+exp(e*(Fvv-
Ehm)/(k*T));
  Plm=Mlx*m0*k*T/(pi*h ^2*dz*10^(-10))*log(1+exp(e*(Fvv-
Elm)/(k*T));
  qc=MT*e^2*h/2/(e0*nr*c*m0^2); % B(hv)/hv/MT2
```

```
72
```

```
rdh=Mrh*m0/2/pi/h_^2/(dz*10^(-10)); % reduce density of
states for C-HH transition
  rdl=Mrl*m0/2/pi/h_^2/(dz*10^(-10)); % reduce density of
states for C-LH transition
  rc=(4*pi*e^2*nr)/(e0*h^2*c^3*m0^2)*MT;% eq.44, p.43 of QWL
  qE sum=dhv.*0;
  qM sum=dhv.*0;
  rh sum=dhv.*0;
  rl sum=dhv.*0;
  for ii=1:1:in ch % C ii to HH ii transition
                                                   % Ee-Ecnz
    fch1=(dhv-Ecm(ii)-Ehm(ii)).*e*Mrh/Me;
in eq.29, p.34 QWL
                                              % Ecnz-Efc
    fch2=(Ecm(ii)-Fcc)*e;
    fch=1./(1+exp((fch1+fch2)./k/T));
                                                   % eq.28,
Fermi-Dirac distribution
    fvh1=(dhv-Ecm(ii)-Ehm(ii)).*e*Mrh/Mhx;
                                                   % Ehnz-Eh
                                              % Efv-Ehnz
    fvh2=(Ehm(ii)-Fvv)*e;
                                                  % (1-fv)
    fvh=1./(1+exp((fvh1+fvh2)./k/T));
eq.28,p.34 QWL
    iic=0;
    for ii1=hv i:hv step:hv_f
      iic=iic+1;
      if fch1(iic) < 0
                                         % normalization
         qnh(iic) = 0;
factor=0 when hv<Eg+Ecnz+Ehnz</pre>
      elseif fch1(iic) == 0
         gnh(iic)=1;
      else
         gnh(iic)=1;
      end
    end
    cos2=(Ecm(ii)+Ehm(ii))*(Mrhz/Mrh)./((dhv-Ecm(ii)-
Ehm(ii))+(Ecm(ii)+Ehm(ii))*Mrhz/Mrh); % cos<sup>2</sup>(Theta_nz)
    sh_{TE=0.5*(1+cos2)};
                                         % transition
strength S TE for HH, p.22 Kenny's thesis
```

```
sh TM=1-\cos 2;
                                % transition strength
S TM for HH, p.22 Kenny's thesis
    gh_{TE}(ii,:) = ((((dhv+Eg).*e). (gc*rdh)).*sh_{TE}).*(fch - 
(1-fvh))*0.01;
    gE sum=gE sum+gh TE(ii,:).*gnh; % C-HH TE gain
    gh TM(ii, :) = ((((dhv+Eg).*e). (gc*rdh)).*sh TM).*(fch -
(1-fvh))*0.01;
    gM sum=gM sum+gh TM(ii,:).*gnh; % C-HH TM gain
                                       % e average for
    sh avg=(2*sh TE+1*sh TM)/3;
C-HH
    rh=((((dhv+Eq).*e*rc).*sh avq).*rdh).*(fch.*fvh); % C-HH
e average spontaneous
    rh sum=rh sum+rh.*gnh; % C-HH e average
spontaneous
  end
  for ii=1:1:in cl % C ii to LH ii transition
    fcl1=(dhv-Ecm(ii)-Elm(ii)).*e*Mrl/Me; % Ee-Ec in
eq.29,p.34 QWL
   fcl2=(Ecm(ii)-Fcc)*e;
                                      % Ec-Efc
   fcl=1./(1+exp((fcl1+fcl2)./k/T));
                                           % eq.28, Fermi-
Dirac distribution
   fvl1=(dhv-Ecm(ii)-Elm(ii)).*e*Mrl/Mlx; % Ev-Eh
   fvl2=(Elm(ii)-Fvv)*e;
                                      % Efv-Ev
   fvl=1./(1+exp((fvl1+fvl2)./k/T)); % (1-fv)
eq.28,p.34 QWL
   iic=0;
   for iil=hv i:hv step:hv f
     iic=iic+1;
```

```
if fcl1(iic) < 0
factor=0 when hv<Eg+Ecnz+Elnz
    gnl(iic)=0;
elseif fcl1(iic) == 0
    gnl(iic)=1;
else
    gnl(iic)=1;
end</pre>
```

```
end
```

```
cos2=(Ecm(ii)+Elm(ii))*(Mrlz/Mrl)./((dhv-Ecm(ii)-
Elm(ii))+(Ecm(ii)+Elm(ii))*Mrlz/Mrl); % cos^2(Theta_nz)
sl_TE=1/6*(5-3*cos2); % transition
strength S_TE for LH
sl_TM=1/3+cos2; % transition strength
S TM for LH
```

% normalization

gl\_TE(ii,:)=((((dhv+Eg).\*e).\(gc\*rdl)).\*sl\_TE).\*(fcl (1-fvl))\*0.01;
 gE\_sum=gE\_sum+gl\_TE(ii,:).\*gnl; % + C-LH TE
gain

```
sl_avg=(2*sl_TE+1*sl_TM)/3; % e average for
C-LH
```

```
rl=((((dhv+Eg).*e*rc).*sl_avg).*rdl).*(fcl.*fvl);
rl_sum=rl_sum+rl.*gnl; % + C-LH e average
spontaneous
```

end

```
% parameters used in the non-Lorentzian lineshape function
  % Broadening factors
  BFv=12+0.3*(N-1.5*10<sup>18</sup>)/(0.2*10<sup>18</sup>); % broadening factor
(meV)
  BFc=8+0.3*(N-1.5*10<sup>18</sup>)/(0.2*10<sup>18</sup>); % broadening factor
(meV)
  BFFv(iN)=BFv;
  BFFc(iN)=BFc;
  tau v=h *1000/BFv/e;
                                   % intraband scattering
time
  tau c=h *1000/BFc/e;
                                   % intraband scattering
time
  t v(iN)=tau v;
  t c(iN) = tau c;
  % effective well widths, modified eq.38, p.112, QWL
  Lc=[1:1:N C]*pi*h ./sqrt(2*Me*m0*Ecm*e); % effective
well width for e, eq.38, p.112, QWL
  Lh=[1:1:N HH]*pi*h ./sqrt(2*Mhz*m0*Ehm*e); % effective
well width for HH, eq.38, p.112, QWL
  Ll=[1:1:N_LH]*pi*h_./sqrt(2*Mlz*m0*Elm*e); % effective
well width for LH, eq.38, p.112, QWL
  % equivalent z component wavevectors, eq.38, p.112, QWL
  kvlt=pi/Lh(1); % equivalent z component wavevector,
kv1_perpendicular, eq.38, p.112, QWL
  kclt=pi/Lc(1); % equivalent z component wavevector,
kc1 perpendicular, eq.38, p.112, QWL
  % minimum of the effective well widths, in eq.41, p.113,
QWL
```

Leh=Lh(1);

Lec=Lc(1);

% inverse screeing length

```
a c=Me*m0*1./(1+exp((Ecm-Fcc)*e/k/T))./Lc; % mcfc(Ecj)/Lcj
   a h=Mhx*m0*1./(1+exp((Ehm-Fvv)*e/k/T))./Lh;
                                                              %
mvfv(Evj)/Lvj for HH
   a l=Mlx*m0*1./(1+exp((Elm-Fvv)*e/k/T))./Ll;
                                                              %
      for LH
  a L=e<sup>2</sup>/pi/h <sup>2</sup>/ee*(sum([a c a h a 1])); % eq.39, p.112,
QWL
  % T(0,kv1t,kv1t), in eq.64, p.124, QWL
  T = 1v = 2/a L + 1/(a L + 4 * kv1t^2);
  T 2v=2/Leh/sqrt(a L)*(1-exp(-Leh*sqrt(a L)));
  T 3v=1/a L-(a L+8*kv1t<sup>2</sup>)/(a L+4*kv1t<sup>2</sup>)<sup>2</sup>;
  T 4v=Leh*kvlt*(Leh*kvlt-pi)*(a L-
4*kvlt<sup>2</sup>)/4/(a L+4*kvlt<sup>2</sup>)<sup>2</sup>/(exp(Leh*sqrt(a L))-1);
  T Lv=(T 1v-T 2v*(T 3v+T 4v))^2; % eq.42, p.113, QWL
  % T(0,kc1t,kc1t), in eq.64, p.124, QWL
  T 1c=2/a L+1/(a L+4*kc1t^2);
  T 2c=2/Lec/sqrt(a L)*(1-exp(-Lec*sqrt(a L)));
  T 3c=1/a L-(a L+8*kc1t<sup>2</sup>)/(a L+4*kc1t<sup>2</sup>)<sup>2</sup>;
  T 4c=Lec*kclt*(Lec*kclt-pi)*(a L-
4*kc1t<sup>2</sup>)/4/(a L+4*kc1t<sup>2</sup>)<sup>2</sup>/(exp(Lec*sqrt(a L))-1);
  T Lc=(T 1c-T 2c*(T 3c+T 4c))<sup>2</sup>; % eq.42, p.113, QWL
  % big K, eq.64, p.124, QWL
  k v=e<sup>4</sup>/(48*pi*ee<sup>2</sup>*h *Lh(1)<sup>2</sup>)*(2*Mhx*m0/h <sup>2</sup>)<sup>2</sup>*k*T;
      % eq.64, p.124, constant in the front
  k c=e<sup>4</sup>/(48*pi*ee<sup>2</sup>*h *Lc(1)<sup>2</sup>)*(2*Me*m0/h <sup>2</sup>)<sup>2</sup>*k*T;
      % eq.64, p.124, constant in the front
  Kv=k v*tau v*T Lv/(1+exp(-Fvv*e/k/T));
                                                                    %
eq.64, p.124, QWL
  Kc=k c*tau c*T Lv/(1+\exp(-Fcc*e/k/T));
                                                                    %
eq.64, p.124, QWL
```

% BGR

```
% n2D 1/m<sup>2</sup>
  N 2D=N*10^{6}*(dz*10^{(-10)});
  d L=0*(-1)*e^2/4/ee/Lh(1)*sqrt(T Lv)*(N 2D);
                                                     %
delta v1, eq.63, p.123, QWL
  % lineshape normalization factor
  G vf=h /2/tau v*(1+exp((Eg-(Eg+dhv1)-
Fvv).*e/k/T))./(1+exp(-Fvv*e/k/T));
  G v1=G vf.*exp((-Kv*e*abs(Eg-(Eg+dhv1)))./k/T);
                                                          %
G v1k11(E), eq.62, p.123, QWL
  G cf=h /2/tau c*(1+exp((Eg-(Eg+dhv1)-
Fcc).*e/k/T))./(1+exp(-Fcc*e/k/T));
  G c1=G cf.*exp((-Kc*e*abs(Eg-(Eg+dhv1)))./k/T);
                                                          Ŷ
G c1k11(E), eq.62, p.123, QWL
  Ln=e*1/pi*(G v1+G c1)./(((Eg-(Eg+dhv1)).*e-
d L).<sup>2+</sup>(G v1+G c1).<sup>2</sup>).*(1-i*((dhv1+Eg-
Eg) + d L/e) . / (G v1+G c1) * e);
  sum L=sum(real(real(Ln)))*hv step;
                                       % normalization
factor
  ic=0;
  chv=Eg-0.08*hv f:hv step:Eg+0.7*hv f;
  for oj=1:length(chv)
    ic=ic+1;
  % non-Lorentzian lineshape
    G vf=h /2/tau v*(1+exp((chv(oj)-(Eg+dhv)-
Fvv).*e/k/T))./(1+exp(-Fvv*e/k/T));
    G v1=G vf.*exp((-Kv*e*abs(chv(oj)-(Eg+dhv)))./k/T);
     % G_v1k11(E), eq.62, p.123, QWL
    G cf=h /2/tau c*(1+exp((chv(oj)-(Eq+dhv)-
Fcc).*e/k/T))./(1+exp(-Fcc*e/k/T));
    G c1=G cf.*exp((-Kc*e*abs(chv(oj)-(Eg+dhv)))./k/T);
     % G_c1k11(E), eq.62, p.123, QWL
```

```
G_L=G_v1+G_c1;
```

```
L=e*1/pi*G L./(((chv(oj)-(Eg+dhv)).*e-
d L).^2+G L.^2).*(1-i*((dhv+Eg-chv(oj))+d_L/e)./G_L*e);
    L=L/sum L;
    gEc(iN,ic) = sum(L.*gE sum)*hv step;
    rch_sum(iN,ic) = sum(L.*rh sum)*hv step;
    rcl sum(iN,ic)=sum(L.*rl sum)*hv step;
           % end of ic
  end
  gEu(iN,:)=gE sum; % unconvolved gain spectra
  Rspu(iN,:)=rh_sum+rl_sum; % unconvolved spontaneous
emission
  [gc max(iN) iMAX(iN)] = max(real(gEc(iN,:)));
  L_cav(iN) = 0.5 \times \log(1/(Ro \times Rb))/(Gama \times gc max(iN) - alfa); 
cavity length
  hv lasing(iN)=chv(iMAX(iN));
                                                     % lasing
energy
  lamda_lasing(iN)=h*c/e/hv lasing(iN);
                                                     %
wavelength
  Rspc(iN,:)=rch_sum(iN,:)+rcl_sum(iN,:);
                                                     %
convolved spontaneous emission
  jrad(iN)=real(sum(Rspc(iN,:))*(hv_step*e)*e/10^4*dz*10^(-
10)); % Jrad in A/(cm<sup>2</sup>)
end % end of N loops
```

```
jth=jrad/eta_i; % Jth in A/cm<sup>2</sup>
```

% plot unconvolved TE gain

%figure(1)

```
%plot(dhv+Eg,gEu)
```

% plot convolved TE gain

%figure(2)

```
%plot(chv,real(gEc))
```

% plot unconvolved spontaneous emission

%figure(3)

```
%plot(dhv+Eg,Rspu)
```

% plot convolved spontaneous emission

%figure(4)

%plot(chv,real(Rspc))

% plots I added

```
%figure(5)
%plot(dhv+Eg,Rspu(1,:),chv,real(Rspc(1,:)))
%xlabel('Energy (eV)')
%title(['Spontanoeus Emission, N=',num2str(Ni)])
%figure(6)
```

```
%plot(dhv+Eg,Rspu(2,:),chv,real(Rspc(2,:)))
```

```
%xlabel('Energy (eV)')
%title(['Spontanoeus Emission, N=',num2str(Ni+N step)])
%figure(7)
%plot(dhv+Eg,Rspu(3,:),chv,real(Rspc(3,:)))
%xlabel('Energy (eV)')
%title(['Spontanoeus Emission, N=',num2str(Nf)])
%figure(8)
%plot(dhv+Eg,gEu(1,:),chv,real(gEc(1,:)))
%xlabel('Energy (eV)')
%ylabel('cm^-1')
%title(['TE Gain, N=',num2str(Ni)])
%figure(9)
%plot(dhv+Eg,gEu(2,:),chv,real(gEc(2,:)))
%xlabel('Energy (eV)')
%ylabel('cm^-1')
%title(['TE Gain, N=',num2str(Ni+N step)])
%figure(10)
plot(dhv+Eg,gEu(3,:),chv,real(gEc(3,:)))
%xlabel('Energy (eV)')
%ylabel('cm^-1')
%title(['TE Gain, N=',num2str(Nf)])
%save 150testx.txt chv -ascii; % me##x.txt: x data for 10<sup>##</sup>
%export=real(Rspc(1,:));
%save 150test1.txt export -ascii; % m#e##.txt: y data for #
x 10<sup>4</sup>#
%export=real(Rspc(2,:));
%save 150test2.txt export -ascii;
%export=real(Rspc(3,:));
%save 150test3.txt export -ascii;
save all2x17x.txt chv -ascii;
export=real(Rspc)/10<sup>49</sup>;
save all2x17.txt export -ascii;
     P InAsSb.m
```

```
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```

function mt=p inassb(T,x)

```
% last modified by Anthony Franz on 7 Aug 97
 % well material for laser B
 % material parameters for InAs(1-x)Sb(x) at temp T K
 % output mt=[Eg Nr gama1 gama2 gama3 Me Mhz Mlz Mhx Mlx S M2 ee]
 % strain effect on energy gap is included
 % matrix elements approximated from k.p model
m0=9.109*10^{(-31)};
e = 1.602 \times 10^{(-19)};
 % Luttinger parameters
 gamalInAs = 20.4; % LB, vol 22a, p 118
 gamalInSb = 36.13; % LB, vol 22a, p 126
gamal = x*gamalInSb + (1 - x)*gamalInAs;
 gama2InAs = 8.3; % LB, vol 22a, p 118
 gama2InSb = 16.24; % LB, vol 22a, p 126
gama2 = x*gama2InSb + (1 - x)*gama2InAs;
 gama3InAs = 9.1; % LB, vol 22a, p 118
 gama3InSb = 17.34; % LB, vol 22a, p 126
gama3 = x*gama3InSb + (1 - x)*gama3InAs;
% energy gap with no strain, from Rogalski, Infrared Phys., vol 29, no
1, p 35, 1989
%egNoStrain=0.411-(3.4*10<sup>(-4)</sup>*T<sup>2</sup>)/(T+210)-0.876*x+0.7*x<sup>2</sup>+(3.4*10<sup>(-</sup>
4) *T*x*(1-x));
% strain correction to energy gap
 a0InAs = 6.0583; % LB1991, p 136
 a0InSb = 6.47937; % LB1991, p 144
a0 = x*a0InSb + (1 - x)*a0InAs; % well lattice parameter in Angstroms
a0s = a0InAs; % substrate lattice parameter
 cllInAs = 8.329; % LB1991, p 137
 cllInSb = 6.669; % LB1991, p 147
cll = x*cllInSb + (1 - x)*cllInAs; % elastic stiffness coefficients in
10<sup>11</sup> dyne/cm<sup>2</sup>
 cl2InAs = 4.526; % LB1991, p 137
 c12InSb = 3.645; % LB1991, p 147
c12 = x*c12InSb + (1 - x)*c12InAs;
 aInAs = -6.0; % Blacha, Phys Stat Sol b, vol126, p 11, table 2
 aInSb = -7.7; % Blacha, Phys Stat Sol b, vol126, p 11, table 2
a = x*aInSb + (1 - x)*aInAs; % hydrostatic deformation potential in eV
bInAs = -1.8; % Blacha, Phys Stat Sol b, vol126, p 11, table 3
bInSb = -2.0; % Blacha, Phys Stat Sol b, vol126, p 11, table 3
b = x*bInSb + (1 - x)*bInAs; % shear deformation potential in eV
es = (a0 - a0s)/a0; % compressive strain (positive) from eq 9, p. 377 of
Zory, QWL
delehh = -2*a*es*(c11 - c12)/c11 + b*es*(c11 + c12)/c11; % change in hh
position relative to cb, eq7,p377,QWL
```

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```

```
delelh = -2*a*es*(c11 - c12)/c11 - b*es*(c11 + 2*c12)/c11; % change in
hh position relative to cb, eq8,p377,QWL
egNoStrain = 0.327479 - (3.79476*10^{(-4)}*T^2)/(T + 346.193);
Eg = egNoStrain + delehh; % strain corrected energy gap in eV
S = delelh - delehh; % strain induced hh-lh gap in eV
% effective masses
 meInAs = 0.0239; % LB1991, p 134
 meInSb = 0.01359; % LB1991, p 142
Me = x*meInSb + (1 - x)*meInAs; % electron effective mass
 mhhzInAs = 0.35; % LB1991, p 134
 mhhzInSb = 0.34; % LB1991, p 142
Mhz = x*mhhzInSb + (1 - x)*mhhzInAs; % z-direction hh effective mass
 mlhzInAs = 0.026; % LB1991, p 134
 mlhzInSb = 0.0158; % LB1991, p 142
Mlz = x*mlhzInSb + (1 - x)*mlhzInAs; % z-direction lh effective mass
 mhhxInAs = 0.35; % LB1991, p 134
 mhhxInSb = 0.42; % LB1991, p 142
Mhx = x*mhhxInSb + (1 - x)*mhhxInAs; % in-plane hh effective mass
 mlhxInAs = 0.026; % LB1991, p 134
 mlhxInSb = 0.0158; % LB1991, p 142
Mlx = x*mlhxInSb + (1 - x)*mlhxInAs; % in-plane lh effective mass
 nInAs = 3.714; % LB1991, p 138
 nInSb = 4.418; % LB1991, p 148
Nr = x*nInSb + (1 - x)*nInAs; % index of refraction
 eInAs = 15.15; % LB1991, p 138
 eInSb = 16.8; % LB1991, p 147
ee = x*eInSb + (1 - x)*eInAs; % static dielectric constant coefficient
 delso = 1.17*x<sup>2</sup> - 0.75*x + 0.39; % spin-orbit splitting in eV, from
Rogalski, 1989
M2 = (1/Me - 1)*((Eg + delso)/(2*(Eg + delso*(2/3))))*Eg*m0*e; %
momentum matrix elements, eq54,p49,QWL
mt=[Eg Nr gama1 gama2 gama3 Me Mhz Mlz Mhx Mlx S M2 ee];
```

#### P\_BarB.m

function mt=p\_barB(T,x,y)
% last modified by Anthony Franz on 7 Aug 97
% barrier material for laser B
% material parameter for In(1-y)Al(y)As(1-x)Sb(x) at temp T K
% output mt=[Eg Nr gama1 gama2 gama3 Me Mh Ml]
m0=9.109\*10^(-31);
e =1.602\*10^(-19);

% final values are interpolated using formula from LB, vol 22a, p 151 % strain in the barrier layer has not been considered % Luttinger parameters gama1InAs = 20.4; % LB, vol 22a, p 118 gama1InSb = 36.13; % LB, vol 22a, p 126 gamalAlAs = 4.04; % P. Lawaetz, Phys Rev B, vol 4, p 3460 gamalAlSb = 4.15; % Lawaetz gamal = (1-x)\*y\*gamalInSb + (1-x)\*(1-y)\*gamalInAs + x\*y\*gamalAlSb + x\*(1-y)\*gama1AlAs; gama2InAs = 8.3; % LB, vol 22a, p 118 gama2InSb = 16.24; % LB, vol 22a, p 126 gama2AlAs = 0.78; % Lawaetz gama2AlSb = 1.01; % Lawaetz gama2 = (1-x)\*y\*gama2InSb + (1-x)\*(1-y)\*gama2InAs + x\*y\*gama2AlSb +x\*(1-y)\*gama2AlAs; gama3InAs = 9.1; % LB, vol 22a, p 118 gama3InSb = 17.34; % LB, vol 22a, p 126 gama3AlAs = 1.57; % Lawaetz gama3AlSb = 1.75; % Lawaetz gama3 = (1-x)\*y\*gama3InSb + (1-x)\*(1-y)\*gama3InAs + x\*y\*gama3AlSb +x\*(1-y)\*gama3AlAs; % energy gaps in eV from Varshni's relation (Physica, vol 34, p 149, 1967)  $alphaInAs = 2.76 \times 10^{(-4)};$ betaInAs = 83;eg0InAs = 0.415; % parameters from Fang, J Appl Phys, vol 67, p 7034, 1990 egInAs = eg0InAs - alphaInAs\*T<sup>2</sup>/(T+betaInAs);  $alphaInSb = 2.7*10^{(-4)};$ betaInSb = 106;eg0InSb = 0.235; % parameters from Fang, J Appl Phys, vol 67, p 7034, 1990 egInSb = eg0InSb - alphaInSb\*T<sup>2</sup>/(T+betaInSb); alphaAlAs = 15.35\*10<sup>(-4)</sup>; % calculated from data from: betaAlAs = 1018;% Monemar, Phys Rev B, vol 8, p 5711, 1973 eg0AlAs = 3.133;% LB vol 22a, p 63 egAlAs = eg0AlAs - alphaAlAs\*T^2/(T+betaAlAs); alphaAlSb = 4.68\*10<sup>(-4)</sup>; % calculated from data from: betaAlSb = 190;% Joullie, Phys Rev B, vol 25, p 7830, 1982 eg0AlSb = 2.384;% LB vol 22a, p 67 egAlSb = eg0AlSb - alphaAlSb\*T<sup>2</sup>/(T+betaAlSb); Eg = (1-x)\*y\*egInSb + (1-x)\*(1-y)\*egInAs + x\*y\*egAlSb + x\*(1-y)\*eqAlAs;% barrier lattice parameter in case I need to add in strain later % a0InAs = 6.0583; % LB1991, p 136 % a0InSb = 6.47937; % LB1991, p 144 % a0AlAs = 5.660; % LB1991, p

```
% a0AlSb = 6.1355; % LB1991, p
    %a0b = (1-x)*y*a0InSb + (1-x)*(1-y)*a0InAs + x*y*a0AlSb + x*(1-
    y) *a0AlAs;
    % effective masses
       meAlSb = 0.259;
       meAlAs = 0.15;
       meInAs = 0.0239; % LB1991, p 134
       meInSb = 0.01359; % LB1991, p 142
   Me = (1-x)*y*meInSb + (1-x)*(1-y)*meInAs + x*y*meAlSb + x*(1-y)*meAlAs;
       mhhzAlSb = 0.336;
       mhhzAlAs = 0.409;
      mhhzInAs = 0.35; % LB1991, p 134
      mhhzInSb = 0.34; % LB1991, p 142
   Mh = (1-x)*y*mhhzInSb + (1-x)*(1-y)*mhhzInAs + x*y*mhhzAlSb + x*(1-x)*(1-y)*mhhzInAs + x*y*mhhzAlSb + x*(1-x)*(1-y)*mhhzAlSb + x*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*(1-x)*
   y) *mhhzAlAs;
      mlhzAlSb = 0.123;
      mlhzAlAs = 0.153;
      mlhzInAs = 0.026; % LB1991, p 134
      mlhzInSb = 0.0158; % LB1991, p 134
  Ml = (1-x)*y*mlhzInSb + (1-x)*(1-y)*mlhzInAs + x*y*mlhzAlSb + x*(1-x)*mlhzAlSb + x*(1-x)*mlhzAlbb + x*(1-x
  y) *mlhzAlAs;
  % index of refraction
     nInAs = 3.714; % LB1991, p 138
     nInSb = 4.418; % LB1991, p 148
     nAlAs = 2.875; % RE Fern, A Onton, J Appl Phys, vol 42, p.3499, 1971
     nAlSb = 3.182; % LB1991, p 85
 Nr = (1-x)*y*nInSb + (1-x)*(1-y)*nInAs + x*y*nAlSb + x*(1-y)*nAlAs;
 mt=[Eg Nr gama1 gama2 gama3 Me Mh Ml];
                          efc.m
 function yf=Efc(N,E,Me,T,d,Ecm)
 8
 ક
 % This subroutine is to calculate the Fermi level of conduction band
 N = carrier density (1/cm^3)
 % Me= conduction band effective mass
T = temperature (K)
% d = active layer thickness (A)
% Ecm= eigenvalue of Ec1,Ec2..
옹
% universal constants
c = 2.998 \times 10^{8};
h = 6.626 \times 10^{(-34)};
h_=h/2/pi;
```

```
k =1.381*10^(-23);
m0=9.109*10^(-31);
e =1.602*10^(-19);
%
c1=Me*m0*k*T/(pi*h_^2*d*10^(-10));
%
ym=c1*log(1+exp( e/k/T* (E-Ecm) ));
yf=sum(ym); % sum of N(subbands)
yf=yf-N*10^6; % [sum of N(subbands)]-[N(given)]. if yf=0, we get the
solution
```

```
efv.m
```

```
function yf=Efv(N,E,Mh,Ml,T,d,Ehm,Elm)
ક
옹
% This subroutine is to calculate the Fermi level of conduction band
N = carrier density (1/cm^3)
% Mh= heavy hole effective mass
% Ml= light hole effective mass
T = temperature (K)
% d = active layer thickness (A)
ş
ક
% universal constant
ዩ
c = 2.998 \times 10^{8};
h = 6.626 \times 10^{(-34)};
h = h/2/pi;
k =1.381*10<sup>(-23)</sup>;
m0=9.109*10^{(-31)};
e = 1.602 \times 10^{(-19)};
옹
ch=Mh*m0*k*T/(pi*h ^2*d*10^(-10));
cl=Ml*m0*k*T/(pi*h_^2*d*10^(-10));
ዩ
yhm=ch*log(1+exp(e/k/T*(E-Ehm)));
ylm=cl*log(1+exp(e/k/T*(E-Elm)));
yf=sum(yhm)+sum(ylm);
yf=yf-N*10^6;
```

# Appendix C: Width Calculations and Convolution of Calculated Spectra

This files convolves the calculated spontaneous emission spectra with the measured pump spectra and creates a file of the results.

```
Off[General::"spell1"]
```

This module calculates the FWHM of a set of x,y data. The x-data should be in one list and the y-data in another list.

```
fwhm[x_, y_] := Module[{max, i, count, xlow, xhigh}, max = Max[y];
For[i = 1; count = 1, y[[i]] <= 0.5 max, count = i; i = i + 1];
xlow = (x[[count+1]] - x[[count]]) (0.5 max - y[[count]])
For[i = Flatten[Position[y, max]][[1]]; count = 1, y[[i]] >= 0.5 max, count = i; i = i + 1];
xhigh = (x[[count+1]] - x[[count]]) (0.5 max - y[[count]])
xhigh = (x[[count+1]] - x[[count]]) (0.5 max - y[[count]])
xhigh - xlow]
```

Read in calculated spectra. Check the file names. The magnitudes have been divided by 10<sup>49</sup>.

```
xData = ReadList["c:\Tony\Thesis\Calculations\SpectralData\unconvx.txt", Number];
yData = ReadList["c:\Tony\Thesis\Calculations\SpectralData\unconv.txt",
Number, RecordLists -> True];
```

```
Length[yData]
Length[yData[[1]]]
```

100



List of the carrier densities associated with each spectrum.

density = Table[i10<sup>16</sup>, {i, 100}];

Find the widths before convolution.

```
calcwidth = {};
Do[dataPoint = fwhm[xData, yData[[i]]];
   AppendTo[calcwidth, dataPoint],
   {i, 1, Length[yData]}]
```

```
Length[calcwidth]
```

100

```
plotData = Transpose[{N[density 10<sup>-17</sup>], N[1000 calcwidth]}];
```

calcPlot =

ListPlot[plotData, PlotRange -> {{0, 10}, {0, 110}}, Axes -> False, Frame -> True, FrameLabel -> {"Density  $(10^{17} \text{ cm}^{-3})$ ", "Width (meV)", "Calculated Widths", ""}]



```
- Graphics -
```

Writes a tab delimited file of width vs. n data called "CalculatvedWidthVsDensity.txt"

```
outputA = OpenWrite["CalculatedWidthVsDensity.txt"]
OutputStream[CalculatedWidthVsDensity.txt, 5]
Do[WriteString[outputA, ToString[plotData[[i, 1]]],
```

```
"\t", ToString[plotData[[i, 2]]], "\n"], {i, 1, Length[plotData]}];
Close[outputA];
```

Load the measured pump spectrum. Subtract the background and normalize it before fitting. The fit is with a  $30^{th}$  order polynomial.

Needs["NumericalMath`PolynomialFit`"]

```
pumpData =
 ReadList["c:\Tony\Thesis\Calculations\DensityVsTime\pumpdata.txt", {Number, Number}];
xpumpData = Table[pumpData[[i, 1]], {i, 1, Length[pumpData]}];
ypumpDataBG = Table[pumpData[[i, 2]], {i, 1, Length[pumpData]}];
ypumpDataBG2 = Table[ypumpDataBG[[i]] - 4000, {i, 1, Length[xpumpData]}];
ypumpData =
 Table[If[ypumpDataBG2[[i]] < 0, 0, ypumpDataBG2[[i]]], {i, 1, Length[xpumpData]}];</pre>
pumpNormalization = Sum[ypumpData[[i]], {i, 1, Length[ypumpData]}];
ypumpData = Table[ypumpData[[i]] / pumpNormalization, {i, 1, Length[xpumpData]}];
pumpFitData = Table[{xpumpData[[i]], ypumpData[[i]]}, {i, 1, Length[xpumpData]}];
Clear[e];
pumpFit = PolynomialFit[pumpFitData, 30];
Length[xpumpData]
1021
xpumpData[[1]]
xpumpData[[-1]]
1.48904
1.60857
dataPlot = ListPlot[pumpFitData, DisplayFunction -> Identity];
fitPlot = Plot[pumpFit[e], {e, 1.489, 1.609}, DisplayFunction -> Identity];
Show[dataPlot, fitPlot, DisplayFunction -> $DisplayFunction, PlotRange -> {0, 0.0005}]
  0.0005
  0.0004
  0.0003
  0.0002
  0.0001
               1.52
                       1.54
                               1.56
                                        1.58
                                                1.6
```

```
- Graphics -
```

Data for pump with same x spacing as the spectra (1 meV).

```
xPump = Table[1.51 + i, {i, 0, 1.6 - 1.51, 0.001}];
yPump = Table[pumpFit[xPump[[i]]], {i, 1, Length[xPump]}];
pumpPlotData = Transpose[{xPump, yPump}];
Length[pumpPlotData]
```

```
91
```



Set up pump with 512 data points for Fourier transform.

```
buffer = Table[0, {i, (512 - Length[xPump] - 1) / 2}];
largeYPump = Flatten[Prepend[yPump, buffer]];
largeYPump = Flatten[Prepend[largeYPump, {0}]];
largeYPump = Flatten[Append[largeYPump, buffer]];
Length[largeYPump]
```

Drop end elements of the spectra so they are 512 elements long.

```
shortYData = Table[Drop[yData[[i]], -(Length[xData] - 512)], {i, Length[yData]}];
Length[shortYData]
Length[shortYData[[1]]]
100
512
```

Set up an x data list with the same spacing between points as the other lists (1 meV).

xConv = Table[0 + 0.001 i, {i, 1, Length[xData]}];

Convolve using Fourier transforms.

```
conv = Table[RotateRight[InverseFourier[Fourier[RotateLeft[shortYData[[i]], 256]]
Fourier[RotateLeft[largeYPump, 256]]], 256],
{i, 1, Length[yData]}];
```



Find the widths of the convolved spectra.

```
convwidth = {};
Do[dataPoint = fwhm[xConv, Abs[conv[[i]]]];
   AppendTo[convwidth, dataPoint],
   {i, 1, Length[yData]}]
Length[convwidth]
```

100

plotConvData = Transpose[{N[density 10<sup>-17</sup>], 1000 convwidth}];
plotConvData[[-1]]

{10., 107.592}

```
convPlot =
ListPlot[plotConvData, PlotRange -> {{0, 10}, {0, 110}}, Axes -> False, Frame -> True,
FrameLabel -> {"Density (10<sup>17</sup> cm<sup>-3</sup>)", "Width (meV)", "Convolved Widths", ""}]
```



- Graphics -

Compare the widths of the calculated spectra with the corresponding convolved spectra.

Show[calcPlot, convPlot, Graphics[Text["Convolved", {0.1, 40}, {-1, -0.1}]],
Graphics[Text["Calculated", {2, 20}, {-1, 0}]],
FrameLabel -> {"Density (10<sup>17</sup> cm<sup>-3</sup>)", "Width (meV)", "Comparison", ""}]



- Graphics -

diff = Table[ convwidth[[i]] - calcwidth[[i]], {i, 1, Length[density]}];
plotDiffData = Transpose[{density 10<sup>-17</sup>, 1000 diff}];

```
ListPlot[plotDiffData, Axes -> False, Frame -> True,
FrameLabel -> {"Density (10<sup>17</sup> cm<sup>-3</sup>)", "(meV)", "Convolved - Calculated", ""}]
```



```
- Graphics -
```

Writes a tab delimited file of width vs. n data called "Fourier512WidthVsDensity.txt"

```
outputB = OpenWrite["Fourier512WidthVsDensity.txt"]
OutputStream[Fourier512WidthVsDensity.txt, 40]
Do[WriteString[outputB, ToString[plotConvData[[i, 1]]],
    "\t", ToString[plotConvData[[i, 2]]], "\n"], {i, 1, Length[yData]}];
Close[outputB];
```

Convolve by summation.

100

```
Length[yConv[[1]]]
```



Find the widths of the convolved spectra.

```
convSumWidth = {};
Do[dataPoint = fwhm[xConv, yConv[[i]]];
   AppendTo[convSumWidth, dataPoint],
   {i, 1, Length[yConv]}]
Length[convSumWidth]
```

100

plotConvSumData = Transpose[{N[density 10<sup>-17</sup>], 1000 convSumWidth}];
plotConvSumData[[-1]]

{10., 107.592}
## convSumPlot =

ListPlot[plotConvSumData, PlotRange -> {{0, 10}, {0, 110}}, Axes -> False, Frame -> True, FrameLabel -> {"Density (10<sup>17</sup> cm<sup>-3</sup>)", "Width (meV)", "Convolved Widths", ""}]



- Graphics -

Compare the widths of the calculated spectra with the corresponding convolved spectra.

Show[calcPlot, convSumPlot, Graphics[Text["Convolved", {0.1, 40}, {-1, -0.1}]],
Graphics[Text["Calculated", {2, 20}, {-1, 0}]],
FrameLabel -> {"Density (10<sup>17</sup> cm<sup>-3</sup>)", "Width (meV)", "Comparison", ""}]



- Graphics -

diff = Table[ convSumWidth[[i]] - calcwidth[[i]], {i, 1, Length[density]}];
plotDiffSumData = Transpose[{density 10<sup>-17</sup>, 1000 diff}];

```
ListPlot[plotDiffSumData, Axes -> False, Frame -> True,
FrameLabel -> {"Density (10<sup>17</sup> cm<sup>-3</sup>)", "(meV)", "Convolved - Calculated", ""}]
```



- Graphics -

Writes a tab delimited file of width vs. n data called "ConvolvedWidthVsDensity.txt"

outputB = OpenWrite["ConvolvedWidthVsDensity.txt"]

OutputStream[ConvolvedWidthVsDensity.txt, 39]

## Appendix D: Carrier Density as a Function of Time

This file maps the carrier density to the delay time using the convolved calculated widths and the measured widths.

```
Off[General::"spell1"]
h = 6.62610<sup>-34</sup>; c = 2.99810<sup>8</sup>; e = 1.60210<sup>-19</sup>;
```

Read in the convolved calculated spectral widths. Check the file name. Format is {density, width} where densities are in units of  $10^{17}$  / cm<sup>3</sup> and widths are in units of meV.

```
calc = ReadList["c:\Tony\Thesis\Calculations\
    DensityVsTime\ConvolvedWidthVsDensity.txt", {Number, Number}];
Length[calc]
100
```

Read in the measured spectral widths. Check the file name. Format is {time, low, high, FWHM} where time is in ps and everything else is in nm.

```
measnm = ReadList["c:\Tony\Thesis\Calculations\DensityVsTime\WidthVsTimenm.txt",
    {Number, Number, Number, Number}];
Length[measnm]
30
```

Convert measured widths to meV and times to ns.

30

Time (ns)	Width (meV)
0.1	81.0064
0.2	77.1179
0.3	74.534
0.4	70.3492
0.5	63.4071
0.6	61.811
0.7	57.7276
0.8	53.9536
0.9	54.9048
1.	52.9698
1.1	50.8694
1.2	53.9536
1.3	47.7102
1.4	47.7102
1.5	44.5611
1.6	46.0618
1.7	42.4294
1.8	43.0309
1.9	42.0887
2.	42.0887
2.1	42.9903
2.2	42.4294
2.3	43.9601
2.4	42.9903
2.5	39.8561
2.6	38.353
2.7	33.7234
2.8	35.2288
2.9	33.7234
3.	33.6704

TableForm[meas, TableHeadings -> {{}, {"Time (ns)", "Width (meV)"}}]

```
ListPlot[meas, Axes -> False, Frame -> True,
FrameLabel -> {"Time (ns)", "Width (meV)", "Measured", ""}]
```



```
- Graphics -
```

This module finds an X value for a Y value equal to A using interpolation on the X,Y list of data.

int[a\_, x\_, y\_] := Module[{i, count, xa}, For[i = 1; count = 1, y[[i]] <= a, count = i; i = i + 1]; xa = x[[count]] - (y[[count + 1]] - a) (x[[count + 1]] - x[[count]]) y[[count + 1]] - y[[count]]

Calculate a table of n vs. t.

xData = Table[calc[[i, 1]], {i, Length[calc]}]; yData = Table[calc[[i, 2]], {i, Length[calc]}]; nt = Table[{meas[[i, 1]], int[meas[[i, 2]], xData, yData]}, {i, Length[meas]}]; Length[nt]

30

Time (ns)	$n (10^{17} \text{ cm}^{-3})$
0.1	7.38127
0.2	7.00538
0.3	6.75402
0.4	6.34424
0.5	5.65386
0.6	5.49289
0.7	5.07474
0.8	4.67895
0.9	4.77991
1.	4.57415
1.1	4.34777
1.2	4.67895
1.3	3.99695
1.4	3.99695
1.5	3.63213
1.6	3.8081
1.7	3.37363
1.8	3.44734
1.9	3.33156
2.	3.33156
2.1	3.44238
2.2	3.37363
2.3	3.56022
2.4	3.44238
2.5	3.04655
2.6	2.84505
2.7	2.15773
2.8	2.39615
2.9	2.15773
3.	2.1492

TableForm[nt, TableHeadings -> {{}, {"Time (ns)", "n (10<sup>17</sup> cm<sup>-3</sup>)"}}]

ListPlot[nt, Axes -> False, Frame -> True, FrameLabel -> {"Time (ns)", "n (10<sup>17</sup> cm<sup>-3</sup>)", "n vs. t", ""}]



- Graphics -

Writes a tab delimited file of carrier density vs. time called "DensityVs.Time.txt"

```
outputA = OpenWrite["DensityVsTime.txt"]
OutputStream[DensityVsTime.txt, 5]
Do[WriteString[outputA,
   ToString[nt[[i, 1]]], "\t", ToString[nt[[i, 2]]], "\n"], {i, 1, Length[nt]}];
Close[outputA];
```

Writes a tab delimited file of meassured width vs. time called "WidthVsTimemeV.txt"

```
outputB = OpenWrite["WidthVsTimemeV.txt"]
OutputStream[WidthVsTimemeV.txt, 6]
Do[WriteString[outputB, ToString[meas[[i, 1]]],
    "\t", ToString[meas[[i, 2]]], "\n"], {i, 1, Length[meas]}];
Close[outputB];
```

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## Vita

Anthony L. Franz was born on 11 July 1970 in Morgantown, WV. He graduated from St. Francis High School in 1988. He studied Space Physics at the U.S. Air Force Academy and earned a Bachelor of Science degree in Physics in 1992. He was commissioned in the U.S. Air Force upon graduation. His first assignment was to the Air Force Technical Applications Center, Patrick AFB, FL, where he served as a Nuclear Systems Evaluator and later as the Advanced Nuclear Detection Program Manager. He was assigned to the Air Force Institute of Technology, Wright-Patterson AFB, OH, in 1996 and received a Master of Science degree in December, 1997. His next assignment was with the Air Force Information Warfare Center, Kelly AFB, TX.

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